

***Fundamentals of Electrocardiography
and Vectorcardiography***

Fundamentals of Electrocardiography and Vectorcardiography

By

LAWRENCE E. LAMB, M.D.

*Director of Cardiology and
Chief, Department of Internal Medicine
Air University
School of Aviation Medicine USAF
Randolph Air Force Base Texas
Consultant in Cardiology
3700th USAF Hospital
Lackland Air Force Base
San Antonio Texas*



CHARLES C. THOMAS PUBLISHER
Springfield Illinois U.S.A.

To My Friend and Teacher
Dr Pierre W Duchosal
Geneva Switzerland

Acknowledgments

I WISH TO TAKE this opportunity to thank the following people who have helped to make this book possible. In particular I wish to thank Colonel Archie A. Hoffman for constructive suggestions, Dr. Julian Ward for calculating the Vectorcardiogram Standardization Charts included in the appendix, and Dr. M. B. Danford and Mr. Richard McNee of the Department of Biometrics for performing the calculations for the Spatial Angle Chart. I am especially pleased with the drawing of the illustrations done by Mr. Leonard F. Carol and assistants. The work done by technicians Sara Johnson, Harvey Hamel, William Yarwood, and Robert Wanner in preparing the electrocardiograms and the vectorcardiogram models was an important contribution. I am indebted to Hedwig Richter and Patricia Dyer for the tedious job of typing the manuscript. The photography was done by Captain Arthur Thiesen, Sergeant Harvey Kohnitz, and staff. To all of these people and the numerous other members of the staff of the School of Aviation Medicine that have helped, I wish to say thank you.

L E L

Contents

	Page
PREFACE	vii
ACKNOWLEDGMENTS	ix
Chapter	
I FUNDAMENTAL VECTOR CONCEPTS	3
II FUNDAMENTAL CELLULAR CONCEPTS	12
The Charged Particle	12
The Doublet	12
The Resting Muscle Fiber	14
Excitation	14
Recovery	15
The Wave Front	15
Summary of Normal Excitation and Recovery	17
Factors Influencing Cellular Recovery	17
Effects of Cell Injury	18
III THE HEART AS A SOURCE OF ELECTRICAL FORCES	19
Atrial Excitation	19
Ventricular Excitation	20
Ventricular Recovery	24
After Potential	26
IV FUNDAMENTALS OF CONDUCTORS	27
Characteristics of Conductors	27
Electrical Field Within a Volume Conductor	27
Infinite Volume Conductors	27
Finite Volume Conductors	28
The Body as a Volume Conductor	28
Location of the Zero Center	29
Remote Electrode and Partial Lead Effect	29
V FUNDAMENTALS OF ELECTROCARDIOGRAPHIC INSTRUMENTS	30
String Gauge Galvanometer	30
The Direct Writing Instrument	31
Cathode Ray Oscilloscope	31
VI FUNDAMENTALS OF ELECTROCARDIOGRAPHIC LEADS	33
Einthoven's Leads	33
Einthoven's Triangle	34
Einthoven's Law	34
The Bipolar Triaxial Reference System	34
Law of Parallelograms	34
The V Lead	34
The Unipolar Triaxial Reference System	34
Hexaxial Reference System	34
Augmented Unipolar Leads	34
Axis by Inspection	34
The Frontal Plane	34

<i>Chapter</i>		<i>Page</i>
	The Transverse Plane	40
	The Twelve Lead Spatial Reference System	42
	Sagittal Plane	44
	Effects of Eccentric Zero Center	44
	Lead Length and Vector Axis	45
	Limitations of Routine Electrocardiography	45
VII	THE NORMAL ELECTROCARDIOGRAM	47
	The P Wave	47
	The PR Interval	47
	The QRS Complex	48
	The Normal ST Segment	55
	The Normal T Wave	55
	The Normal U Wave	56
	The Normal QT Interval	56
	The Mean Vector	56
	The Mean of QRS Vector	57
	Normal Mean QRS Axis	60
	The Mean T Vector	60
	The Mean Spatial QRS T Angle	61
VIII	FUNDAMENTALS OF VECTORCARDIOGRAPHY	66
	The VCG System	68
	Index of Maximum Potential and Potention Seconds	71
	The Normal Vectorcardiogram	72
IX	CARDIAC ENLARGEMENT	82
	Atrial Enlargement	82
	Left Ventricular Enlargement	82
	Right Ventricular Enlargement	84
X	CONDUCTION DEFECTS	91
	Left Bundle Branch Block	91
	Right Bundle Branch Block	93
	The S ₁ S S ₃ Pattern	94
	Intra Ventricular Conduction Defects	96
	Accelerated Conduction	96
XI	PERICARDITIS	105
XII	MYOCARDIAL INFARCTION AND ARTERIOSCLEROTIC HEART DISEASE	108
	Myocardial Infarction	108
	Arteriosclerotic Heart Disease	116
XIII	DRUGS AND METABOLISM	117
	Digitalis	117
	Quinidine	118
	Hypopotassemia	118
	Hyperpotassemia	121
	Calcium	121
	Thyroxin	121
XIV	ARRHYTHMIAS	122
	Sinus Rhythms	122

Contents

<i>Chapter</i>	<i>Page</i>
Atrial Rhythm	124
Nodal Rhythms	126
Ventricular Rhythms	129
AV Block	130
\\ THE INTERPRETATION	132
INDEX	139

<i>Chapter</i>		<i>Page</i>
	The Transverse Plane	40
	The Twelth Lead Spatial Reference System	42
	Sagittal Plane	44
	Effects of Eccentric Zero Center	44
	Lead Length and Vector Axis	45
	Limitations of Routine Electrocardiography	45
VII	THE NORMAL ELECTROCARDIOGRAM	47
	The P Wave	47
	The PR Interval	47
	The QRS Complex	48
	The Normal ST Segment	55
	The Normal T Wave	55
	The Normal U Wave	56
	The Normal QT Interval	56
	The Mean Vector	56
	The Mean QRS Vector	57
	Normal Mean QRS Axis	60
	The Mean T Vector	60
	The Mean Spatial QRS T Angle	61
VIII	FUNDAMENTALS OF VECTORCARDIOGRAPHY	66
	The VCG System	68
	Index of Maximum Potential and Potention Seconds	71
	The Normal Vectorcardiogram	72
IX	CARDIAC ENLARGEMENT	82
	Atrial Enlargement	82
	Left Ventricular Enlargement	82
	Right Ventricular Enlargement	84
X	CONDUCTION DEFECTS	91
	Left Bundle Branch Block	91
	Right Bundle Branch Block	93
	The S ₁ S S ₃ Pattern	94
	Intra Ventricular Conduction Defects	96
	Accelerated Conduction	96
XI	PERICARDITIS	105
XII	MYOCARDIAL INFARCTION AND ARTERIOSCLEROTIC HEART DISEASE	108
	Myocardial Infarction	108
	Arteriosclerotic Heart Disease	116
XIII	DRUGS AND METABOLISM	117
	Digitalis	117
	Quinidine	116
	Hypopotassemia	118
	Hyperpotassemia	121
	Calcium	121
	Thyroxin	121
XIV	ARRHYTHMIAS	122
	Sinus Rhythms	122

Contents

<i>Chapter</i>	<i>Page</i>
Atrial Rhythm	124
Nodal Rhythms	126
Ventricular Rhythms	129
AV Block	130
VII THE INTERPRETATION	132
INDEX	139

***Fundamentals of Electrocardiography
and Vectorcardiography***

Fundamental Vector Concepts

ELECTROCARDIOGRAPHY strives to measure potential (forces) created by the heart in terms of magnitude direction and duration of action. Vector concepts greatly simplify analysis of these forces. The use of vectors to express mathematical quantities is as old as mathematics itself. They were applied to electrocardiography by Einthoven. The understanding of electrocardiography begins with an understanding of simple vector principles.

What Is a Vector? It is a symbol used to describe the characteristics of a force. The magnitude of the force is represented by the length of the vector. The arrow head (terminus) of the vector indicates the direction of the force. A vector used to describe a force in two dimensions (flat surface) is drawn as a flat arrow. A spatial vector representation is used to describe a three dimensional force (Fig 1). The duration of action of a force is not expressed by a

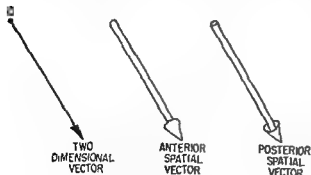


Figure 1

vector unless the magnitude has been converted to time units. A force of one dyne acting for ten sec is still a one dyne force represented by one unit length (1 unit equals 1 dyne). Expressed

in time units is $\text{dynes} \times \text{seconds}$ such a force would be equal to ten dyne seconds and represented by ten units length (1 unit equals 1 dyne second).

What Is the Point of Origin? The maximum effect of a force occurs at a point. The force is said to act on this point. In vector terminology this point is called the point of origin. One end of any vector is its point of origin and the other end is the head (terminus) indicating the direction of the vector and extent of its magnitude.

What Is an Instantaneous Vector? A vector without duration of time is an instantaneous vector. It acts only at a point in time. In electrocardiography it is often of interest to speak of a vector acting at one particular time interval; thus the vector acting momentarily at 0.4 sec after the onset of an event is spoken of as the 0.4 sec vector. Such a vector is an instantaneous vector and acts only at that time interval. At 0.5 sec after the onset of the same event the vector might be entirely different in magnitude and direction. The 0.5 vector would be another instantaneous vector.

What Is a Resultant Vector? When two or more vectors are acting on a common point of origin their net effect can be represented by a single vector. Such a vector is called a resultant vector (Fig 2). It represents the effective force of all the vectors. The resultant vector is not equal to the magnitude of all the vectors acting on the point nor does it have a consistent relationship to their magnitude.

What Is a Coordinate? Any straight line drawn through the point of origin is a coordi-

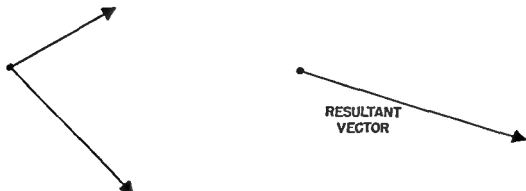


Figure 2

nate Its length can be marked off into units of measure The effect of a vector in the direction defined by the coordinate (line) can be determined by constructing a perpendicular from the coordinate to the tip of the vector (Fig 3) The units along the coordinate equals the effect of

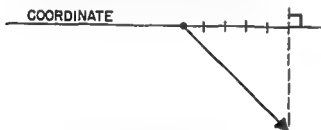


Figure 3

the vector in that direction This value is the coordinate value of the vector The value may be expressed as positive or minus units by designating one end of the coordinate positive and the other negative

The coordinate value of the vector is spoken of as its projection upon the coordinate All electrocardiographic leads are coordinates and measure forces that are projected upon them Any vector can be described in terms of its projection upon three mutually perpendicular coordinates (X Y Z) The X coordinate (transverse) defines the left to right location of the terminus The Y coordinate (vertical) defines the vector terminus above or below the point of origin The Z coordinate (sagittal) locates the vector terminus anterior or posterior to the point of origin (Fig 4)

What Are Component Vectors? The units of magnitude measured upon the coordinates X Y

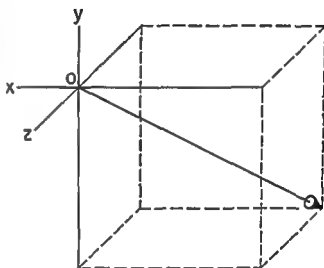


Figure 4

Z can be expressed as vectors The three vectors X Y Z are component vectors The resultant of three such vector components is equal to the magnitude and direction of the spatial vector They are perpendicular vector components (Fig 5) Any spatial vector can be resolved into its three perpendicular components The magnitude of the spatial vector can be calculated from its perpendicular components from the simple formula

$$(\text{Spatial vector})^2 = X^2 + Y^2 + Z^2$$

Any spatial vector may have multiple component vectors and multiple coordinates but it can have only three mutually perpendicular coordinates and three mutually perpendicular components

How Is the Coordinate Value of a Vector Calculated? When the angle between a vector and a given coordinate is known the projection

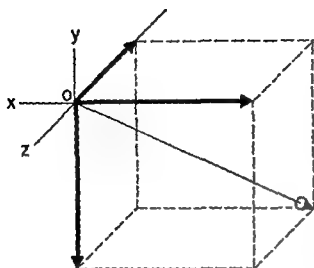


Figure 5

of the vector on the coordinate can be calculated without constructing a perpendicular (Fig 6). This is done by using the principle of the right triangle. The vector is the hypotenuse of a right triangle and the coordinate is the adjacent side. The cosine* of the angle times the

The cosine of an angle is equal to adjacent side divided by the hypotenuse.

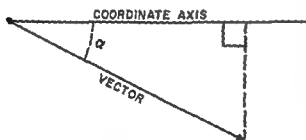


Figure 6

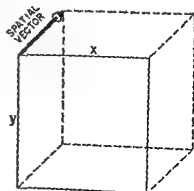
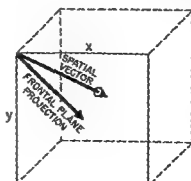
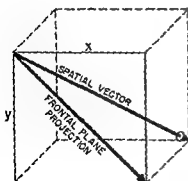


Figure 7

magnitude of the vector equals its coordinate value

$$\cos \alpha \times \text{Vector Magnitude} = \text{Coordinate Value}$$

What Is a Plane? A plane expresses two dimensions of a vector and has two perpendicular coordinates. It is a flat surface. The *frontal plane* (X Y) is made up of the X Y coordinates. The *transverse plane* (X Z) is composed of the X Z coordinates. The *sagittal plane* (Z Y) is the Z Y coordinates. These planes are mutually perpendicular to each other. Any two of them can be used to determine a spatial vector as two perpendicular planes include all three mutually perpendicular coordinates (X Y Z).

The principle of measuring a vector from its projection upon a plane or flat surface is frequently used in electrocardiography. The magnitude of a vector measured by a plane diminishes as the vector is rotated away from its flat surface (Fig 7). Considering the frontal plane as the vector is rotated away from the X Y coordinates the frontal plane vector becomes smaller. Finally, when the vector is parallel to the Z coordinate its entire magnitude is measured by the Z coordinate. Such a vector has no projection upon the X or Y coordinate and has no magnitude value in the frontal plane. Both X and Y are perpendicular to Z and the frontal plane is perpendicular to such a vector. A simple rule is illustrated: there is no force at any point on a plane perpendicular to a vector. The direction of a vector can be determined by locating its perpendicular plane as the vector is 90° away from this plane.

What Is the Law of Parallelograms? The resultant of two vectors acting upon a common point can be determined by constructing a parallelogram (Fig 8). The two vectors are two adjacent sides (a , b) of the parallelogram. The diagonal of the parallelogram is the resultant vector (c). The magnitude of the vectors act

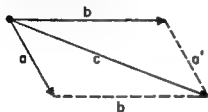
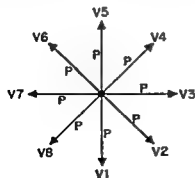


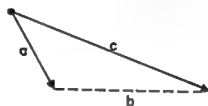
Figure 8

ing on the common point can be determined from the sides of the parallelogram (a , b). Note that $b = b'$ as they are opposite sides of the parallelogram. Knowing only a and c , the magnitude of the vector b can be determined by merely measuring the distance between vector a and c or the length of b . The two adjacent sides of the parallelogram equals the magnitude of the acting vectors and the diagonal equals their effect or resultant.

It is very important to realize that the diagonal of the parallelogram is not equivalent to the magnitude of the acting vectors. Any triangle is a semiparallelogram. When one side of a triangle is a component (acting) vector (a) and the other the resultant vector (c), the other component vector (b) can be determined from the law of parallelograms.



The Law of Simple Consecutive Vector Addition The addition of vectors upon a common point one after another will cause the resultant vector to change with the addition of each vector. Consider eight vectors of equal magnitude (P) consecutively added to each other upon a common point (O) and directed 45° away from



each other (Fig 9). The addition of vector 2 to vector 1 creates the resultant 1. Note that R_1 is the diagonal of a semiparallelogram while V_1 and V_2 are the adjacent sides of the parallelogram. The addition of vector 3 to vectors 1 and 2 creates the resultant vector 2.

The addition of each new vector causes the resultant vector to rotate a distance equal to the added vector's magnitude. The rotating resultant vector describes an external pathway comprised of the sides of a polygon. The length of the sides of the polygon is equal to the total magnitude of all the vectors acting on the common point. In this instance the magnitude of the eight vectors acting on the common point is $8P$. The sum of the length of the sides of the polygon is also $8P$. The graph of the rotating resultant vectors is a polygon. The resultants

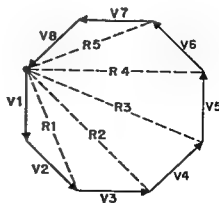


Figure 9

originating from the center of origin may be called central resultants. Note that the magnitude of the acting vectors is never equal to the central resultants. A law may be formulated: *The consecutive addition of vectors upon a common point will cause the central resultant to rotate describing a polygon. The length of the sides of the polygon is equal to the magnitude of the vectors added to each other.*

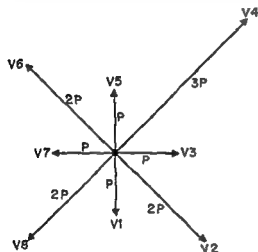
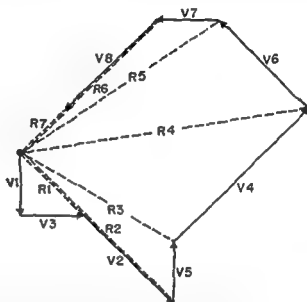


Figure 9 The irregular shaped polygon is created by unequal vectors acting upon the point of origin. Note that the simultaneous action of such unequal vectors would have a resultant in this case R_7 . The resultant of several vectors acting on a common point can always be determined in this manner. In the previous illustrations the resultant has been zero and the polygon was a closed polygon.

The Law of Simple Consecutive Vector Subtraction The subtraction of vector forces one after another from a common point creates a similar situation as consecutive addition. Given eight vectors of equal magnitude (P) directed

Figure 9 is drawn to scale as geometric proof of the law. The method of polygon formation must apply to every situation wherein vectors are assumed to act upon a common point or center. The polygon finds its origin in the law of parallelograms. Whenever instantaneous resultant vectors are assumed to act on a point the sides of a polygon can be constructed. The sides of the polygon then represent the magnitude of the component vectors. The reader is referred to White Harvey E. *Classical and Modern Physics* New York Van Nostrand Company Inc. 1940 pp. 25-6 and Kimball A. L. *A College Text Book of Physics* Fifth Edition New York Henry Holt and Company 1939 p. 11.

45 $^{\circ}$ from each other there will be no effective force until subtraction begins. With the subtraction of vector 1 an effective force is created by the remaining seven vectors. The subtraction of vector 2 changes the resultant again. The resultant vector changes with each addition or subtraction. A law may be formulated: *The successive subtraction of vectors from a common point causes a resultant vector to be*



rotated describing a polygon. The length of the sides of the polygon is equal to the magnitude of the vectors subtracted. In either addition or subtraction the shape of the polygon will also depend upon the order of addition or subtraction (Fig. 10).

The Law of Multiple Simultaneous Vector Addition The addition of vectors upon a common point may be complicated by the simultaneous addition of more than one vector. Consider two vectors a and b added simultaneously to a previously existing vector A (Fig. 11). Vectors a and b have a resultant effect c the diagonal of a parallelogram. The action of these two vectors is equal to vector c . When vectors

a and b are added to vector A it is exactly the same as if vector c had been added to vector A knowing vector A and the new resultant (B) created by the addition of vector c one can determine vector c by measuring the distance between A and B . A law may be formulated: *The simultaneous addition of two or more vectors to a previously existing vector will cause it to rotate describing one side of a polygon. The length of the side is equal to the magnitude of the resultant of the simultaneously added vectors.* A similar relationship exists for subtraction.

What Is a Coordinate Graph? Electrocardiographic leads are coordinate graphs in contrast

to vectorcardiograms which are polygons. They are vectors graphed upon a coordinate in a linear fashion. Extend coordinates X and Y through the circular type graph of consecutive vector addition (Fig. 12). Each central resultant can be projected upon the coordinates in a linear fashion. This creates a coordinate graph of the x components of central resultants and another graph of the y components of the central resultants. Note that such graphs are graphs of the central resultants and not the sides of the polygon. As shown above the magnitude of vectors acting upon a common point

is measured by the sides of a polygon. By indirect calculation or geometric construction the polygon can be obtained from linear coordinate graphs but such a procedure must be carried out before one can speak correctly in terms of vector magnitude.

What Is an Absolute Linear Graph? The central resultants may be graphed upon a time base to form a linear graph (Fig. 13). A popular application to vectorcardiography is to so graph the instantaneous resultant spatial vectors. It is clear from a fundamental vector approach that such central resultants in no way represent the magnitude of the component vectors acting upon a common point and unless one knows the

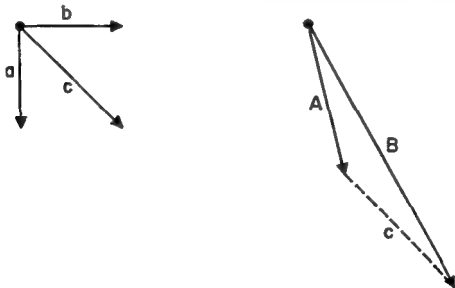


Figure 11

distinction to vectorcardiograms which are polygons. They are vectors graphed upon a coordinate in a linear fashion. Extend coordinates X and Y through the circular type graph of consecutive vector addition (Fig. 12). Each central resultant can be projected upon the coordinates in a linear fashion. This creates a coordinate graph of the x components of central resultants and another graph of the y components of the central resultants. Note that such graphs are graphs of the central resultants and not the sides of the polygon. As shown above the magnitude of vectors acting upon a common point

is measured by the sides of a polygon. By indirect calculation or geometric construction the polygon can be obtained from linear coordinate graphs but such a procedure must be carried out before one can speak correctly in terms of vector magnitude.

angles between such central resultants the sides of the polygon cannot be reconstructed. These measurements have been called the absolute vectorcardiogram. A linear graph of the central resultants is called an absolute graph. Actually the use of the term absolute is a misnomer in

Each side of the polygon can be determined by obtaining its x and y component. The x component is the difference in amplitude of the central resultant projection upon x coordinate for that time interval and the y component is the difference in amplitude of the central resultant projection upon the y coordinate. The side for this time interval or for the added vector is then obtained from the formula $x^2 + y^2 = \text{Side}$ or by simple graphic construction.

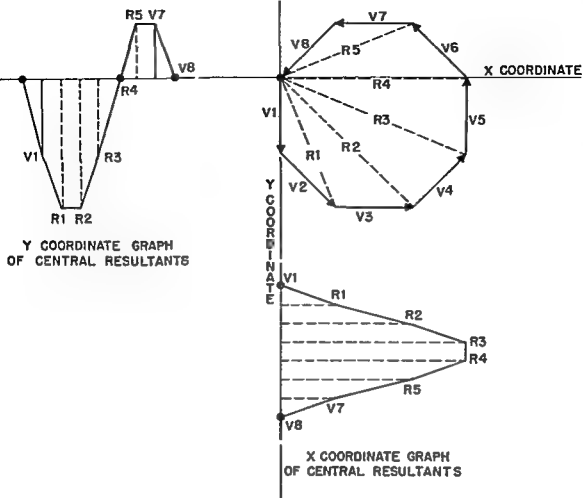


Figure 12

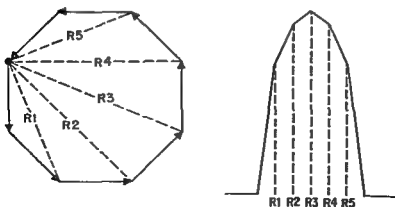


Figure 13

that resultant vectors are not absolute measurements and such measurements of absolute quantities must be approached from the polygon principle.

Two different polygons can be constructed with entirely different length of sides and identical central resultants (Fig 14). Even

from the same point (o) creating a new resultant (b). The magnitude of the force subtracted is equal to the distance between the two resultants (a and b). If this distance is equivalent to a force of two dynes and its duration of action was five seconds it can be expressed as ten dyne seconds ($2 \text{ dynes} \times 5 \text{ sec}$).

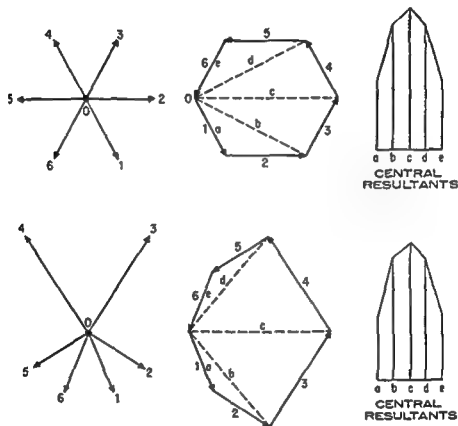
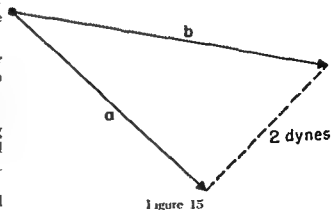


Figure 14

though the central resultants are the same the component vectors are not at all the same. This simple illustration demonstrates the fallacy of using instantaneous resultant vectors to measure magnitude of component vectors.

Expression of Vector Magnitude In Time Units Once the magnitude of a force is known it can be converted to units of time by multiplying its magnitude by its duration of action. When we are concerned with forces acting about a point the magnitude factor is obtained from the sides of the polygon. Consider a force of three dynes (a) acting upon a point (Fig 15). Five seconds later a force is subtracted

The Mean Vector from the Area of Ordinate Graphs A popular concept in electro



cardiography is to obtain the "mean vector" (mean QRS or T) from calculations based on coordinate graphs (the leads themselves). The first step is to obtain the triangular area enclosed on a given coordinate. This is the coordinate value for the mean vector. Another coordinate value is then obtained in a similar manner. A perpendicular is then constructed

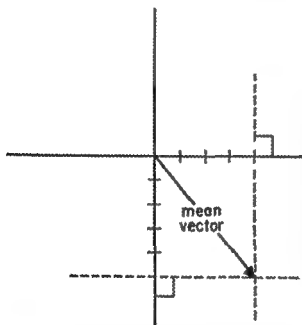


Figure 16

to each of the two coordinates marking off their unit values. The point of intersection between the two perpendiculars is the terminus or the "mean" vector. The magnitude of the vector is depicted by its length and its direction by its axis and terminus (Fig. 16).

The base of the triangle is a function of time and the amplitude of the complex is a function of magnitude. The height times the base divided by two is the triangular area. In the event that a triangular area exists above the coordinate it is positive and if below it is negative. If part of the coordinate graph has a positive value and the other part negative the two triangular areas are subtracted from each other or added by algebraic addition. This value represents the mean value measured by such a coordinate.

The term "mean" vector as determined by the above method is a misnomer. The vector magnitude has no consistent relationship to the magnitude of vectors acting about a common point. The basic premise for magnitude is in error as magnitude must first be determined from the polygon principle and then converted to time units. Nevertheless such graphs are satisfactory for determining the direction of the resultant vector created by multiple forces acting about a point.

Instantaneous Vector Versus Polygon Sides
A simple graphic demonstration of the difference in values obtained from instantaneous resultant vectors and the polygon is exemplified by travel. Driving by car from San Antonio to Houston to Dallas one describes two sides of a polygon. The actual road mileage is the sides of the polygon and can be equated to the work performance of the auto. The air mileage from San Antonio to Dallas is the resultant or instantaneous vector being the resultant of the sides of the polygon. Obviously one cannot equate work done by the auto to the air mileage nor can any function derived from the air mileage alone be expected to represent the auto's true activity (Fig. 17).

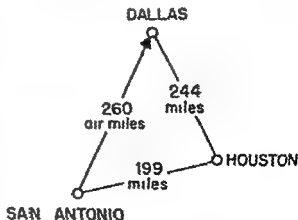


Figure 17

II

Fundamental Cellular Concepts

THE ELECTRICAL EVENTS of the heart conform to basic principles of electrical physics. The events concern themselves with particles containing electrical energy. This in turn is related to the atomic structure of the elements or electrolytes found in the tissue mass and surrounding media. An intelligent understanding of the cell is a source of electrical activity requires at least a handshaking acquaintance with the fundamental behavior of particles bearing electrical energy.

THE CHARGED PARTICLE

Any particle containing electrical energy is a charged particle. Under proper environmental conditions this particle exerts a force. The magnitude of the force created by a particle is called its potential*. All charged particles are

Potential is correctly defined as work. A unit of potential one volt is equal to 1/300 electrostatic unit or 1/300 stat volt. An electrostatic unit difference in potential between two points exists when one erg of work is done to move one stat coulomb of positive charge from the point of least potential to the point of more positive potential. One erg is one dyne cm. or a force of one dyne acting a distance of one cm. Thus potential difference can be expressed in dyne cm. A dyne is a fundamental unit of force and is correctly given the dimension of magnitude.

Potential at a point is also expressed as work. The difference in potential is expressed between the point in question and earth potential, the latter being designated as zero potential.

Potential depends upon the charge that a particle bears but is an expression of force not charge. Charge refers to the number of electrons contained by the particle. Current expressed in amperes is the amount of charge (coulombs) flowing per second. For further detail the reader is referred to Gilbert Norman E. *Electricity and Magnetism* Third Edition New York The Macmillan Company 1930.

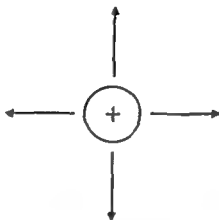


Figure 18

surrounded by an electrical field. The field is created by the transmission of potential into the surrounding media. Any point in this field has a potential value. The potential decreases at greater distances from the particle in accordance with the characteristics of the surrounding media. Since its potential is transmitted in all directions the particle has no effective electrical force i.e. the resultant force is zero (Fig 18).

THE DOUBLET

An effective electrical force or resultant force is created when two charged particles of different potential values are close enough to each other for overlapping of their electrical fields (Fig 19). The force has direction and magnitude. The magnitude is equal to the difference in potential between the two particles. Potential difference is expressed in practical units as voltage. By convention the particle of lesser charge is called negative and the particle of

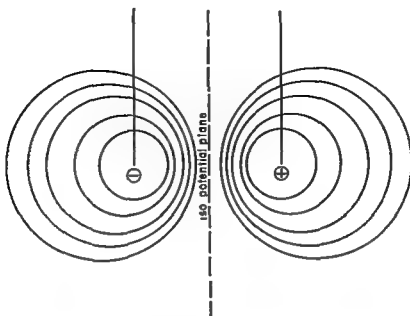


Figure 19

greater charge positive. The electrical field surrounding the particle of greatest potential is called the positive electrical field and the field surrounding the point of least potential is called the negative electrical field. Between the two particles a zone must exist which is as positive to the negative particle as it is negative to the positive particle. This zone is created by the overlapping of the two electrical fields. The forces emitted from each particle tend to neutralize each other in the area between the two particles. This plane of neutralization is called the *isoelectric* or *isopotential plane*.

On the outer side of the positive particle the potential forces are unopposed and of course larger than those on the outer side of the negative particle. This difference in potential value creates a force directed away from the isopotential plane and toward the positive field. The maximum line of force must be in line with the axis between the positive and negative particles.

*A particle may have less than zero potential in relation to earth potential. A particle may have negative potential in relation to another particle. As an illustration if one particle has 35 units of potential and the other is 5 units, the potential difference is 30 units. It is of no consequence where the zero point is placed. One may express their relative values as -15 and +15 with the same mathematical relationship.

Such a line is perpendicular to the isopotential plane. The force acts as if it acted on a point where the line of force intersects the isopotential plane. This point is called the *zero center*. It should be noted that this does not mean there is no potential at that point. It is a point of electrical neutrality. The force created by the electrical fields behaves as if it acted upon this point. The term *"zero center"* is an electrical term.

Two particles of different potential as described above create a force which may be described as a vector (Fig. 20). Its point of origin

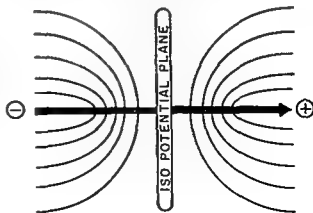


Figure 20

the zero center and its direction is toward the positive electrical field. The vector magnitude depends upon the difference in potential between the particles, their distance apart, and the character of the surrounding media. Two such particles are called a *doublet*. The force created by the doublet is called the *doublet vector*.

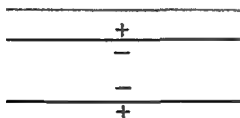
THE RESTING MUSCLE FIBER

The electrolytic content of the normal resting muscle fiber differs from the electrolytic content of its normal external environment. Consequently, there is a continuous diffusion of ions across the cell membrane. There is a natural tendency for ions to migrate from a location of higher density to one of lower density, thereby reaching a state of equilibrium. The difference in ionic concentration on either side of the membrane is the driving force of migration. The ionic activity may be altered by environmental factors. The diffusion of ions across the cell membrane also depends upon the character of

(Fig. 21) A positive electrical field surrounds the muscle fiber. Thus, forces are directed externally at all points across the cell membrane. It is as if electrical doublets existed at all points around the cell. The resultant force is zero and the resting cell creates no effective force in its surrounding media. However, a force does exist across the cell membrane because the internal environment is negative with respect to the external environment. This potential difference is the *resting membrane action potential*. Experimental measurements have shown it to have a value of 50 to 90 mv.

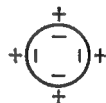
EXCITATION

When the muscle fiber is stimulated, a change occurs in the cell membrane. Stimulation may be accomplished by mechanical, chemical, or electrical means. The membrane potential is altered and its behavior as a filter is changed. This permits a change in dynamic diffusion of electrolytes across the membrane. The initial event is the migration of sodium ions from the external



LONGITUDINAL SECTION

RESTING MUSCLE FIBER



CROSS SECTION

Figure 21

the membrane itself. In this sense the membrane acts as a filter (dielectric constant) regulating the rate of diffusion between the two areas of different ionic content.

The diffusion of ions across the membrane and the character of the membrane filter are relatively stable in the normal resting state, thus creating a state of dynamic equilibrium. The external environment of the cell is relatively positive with respect to the internal environment

environment to the internal environment, or an inward current of sodium. Stimulation increases the permeability of the cell membrane to sodium and since there is a higher concentration of sodium externally, sodium migrates inward.

The change in the potential of the membrane and the change in electrolytic content of the internal and external cellular environment creates a positive electrical field within the cell.

with respect to its external environment (negative field). One may call this a reversal of the membrane action potential. The area of the muscle fiber which has undergone this change is said to be in the *excited state*. The force created by the difference in potential is directed inward from all external points perpendicular to the cell membrane. The resultant force is zero and there is no effective force in the external media distant to the cell. A rule may be formulated: *An area of muscle in the excited state creates no effective force in its surrounding media.*

During excitation of a muscle fiber there must exist an area in the excited state and an area in the resting state. At the junction of these two areas the active process of excitation is occurring. This boundary is the *wave of excitation* (Fig. 22). As pointed out above, neither the

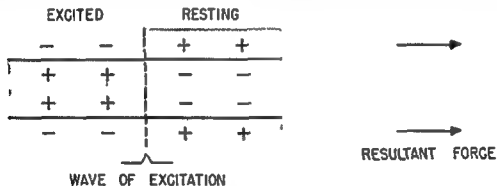


Figure 22

area in the excited nor resting state creates a force in the surrounding media. The force created at any instant in the external media is due to the boundary between muscle areas in different electrical states in this case the wave of excitation. Such a force is created at the surface of the cell and not across the cell membrane.

As the wave of excitation progresses along the fiber it forms a boundary at any one instant between recovered (resting) tissue and excited tissue (Fig. 23). This boundary is the *wave of recovery*. It has exactly the same characteristics as the wave of excitation. A *wave of recovery* creates a force in the surrounding media directed toward the area of resting (recovered) tissue and away from excited tissue.

THE WAVE FRONT

When a wave front is a flat surface the force it creates is directly proportional to its area. Although the magnitude may be altered by

The force parallel to the surface may be explained by considering the field created perpendicular to flowing current as in the Oersted experiment. The wave front defines the border between current flowing in two different directions. The bidirectional current creates a force perpendicular to the direction of current flow and perpendicular to the wave front.

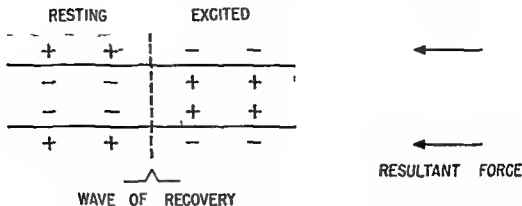


Figure 23

changes in electrolytic and physical environment (the density of doublet charge) the relationship between wave area and magnitude persists. Thus the larger the flat surface area of the wave front the larger the magnitude of the force (potential). The wave front may be regarded as having forces acting perpendicular to its surface at all points.

If the magnitude of the force created by the wave front is known in mv and its duration of action in sec is determined its value can be expressed in $\text{mv} \times \text{sec}$. A force of 10 mv across a wave front lasting 10 sec is expressed as 100 mv sec. Expression of potential differences acting through a period of time in this manner is frequently used in electrocardiography.

The wave front is not always flat. When it is not flat the force created in the surrounding medium is no longer proportional to its surface area. Vector doublets will not be perpendicular at all surface points. By vector addition when forces are not acting parallel and in the same direction the resultant of their action is not equivalent to the sum of their individual magnitude. The effective force of a wave front is equivalent to the resultant of the doublet vectors acting perpendicular to its surface.

Given a conical wave front one can calculate its resultant vector (Fig 24). Since all the doublet vectors are acting perpendicular to the wave front their resultant will be perpendicular to the base (assuming even distribution of doublet charges). This exemplifies an important principle of wave fronts: *whenever a wave front*

bounds a single flat surface area its resultant will be perpendicular to the flat surface. The magnitude of the resultant vector will be directly proportional to the area of the base. *Whenever a single flat surface area is bounded by a wave front the magnitude of its resultant vector is directly proportional to the area of the flat surface.*

A wave front may have more than one flat surface area. A cone with $1/6$ of its arc cut away

This simple principle is verified by the theorem of Gauss. In part there are two essential equations. The electrical intensity at a point in field due to multiple distributions of electrical charge is found in the equation

$$\iint E \cos \theta \, ds = \frac{4\pi}{k} Q$$

The intensity (E) at a point on the surface distant to a field created by a uniformly charged spherical shell is

$$E = \frac{Q}{k r^2} \text{ dynes per stat coulomb}$$

Q is the charge, k is the dielectric constant of the surrounding medium and r is the distance between the surface point and the center of the spherical shell. It should be noted that effect of the spherical shell is as if the intensity originated from the center of the sphere. For further detail the reader is referred to Gilbert Norman E. Electricity and Magnetism Third Edition New York The Macmillan Company 1950.

An equation by derivation from the work of Gauss was used by Wilson to calculate point potential at a distance

from the source $V = \frac{\phi S \cos \theta}{r}$ where V was the point

potential ϕ the density of charge S the flat surface area θ the angle between the axis of the flat disk and a line drawn to the potential point and r the distance between the point from the flat disk. Wilson F. N. MacLeod A. G. and Barker P. S. *The Distribution of the Currents of Action and Injury Displayed by Heart Muscle and other Excitable Tissues* Univ. of Mich. Studies Scientific Series Ann Arbor Univ. of Mich. Press 1933 Vol 18 p 58.

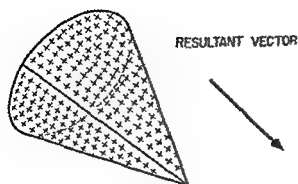


Figure 24

has two flat surface areas (Fig 25). The resultant of the wave front is equal to the resultant of two vectors: one representing the base of the cone and the other flat surface area created by the 1st conical area.

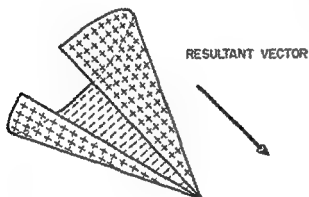


Figure 25

SUMMARY OF NORMAL EXCITATION AND RECOVERY

The following simple rules may be formulated to summarize excitation and recovery of a normal muscle fiber:

- 1) Muscle fibers in the resting state create no effective external force.
- 2) Muscle fibers in the excited state create no effective external force.
- 3) The boundary between muscle areas in different electrical states (excited and resting) is a wave front and creates a force directed toward the area of muscle in the resting or recovered state.
- 4) The magnitude of the force created by a

wave front is the resultant of the doublet vectors acting everywhere perpendicular to the wave front.

5) Under normal circumstances recovery begins at the same site as the onset of excitation. This creates a force opposite the force of excitation (rule 3).

FACTORS INFLUENCING CELLULAR RECOVERY

Recovery of the isolated muscle strip is very sensitive. Numerous factors may alter its usual process. Warming speeds up recovery. If one end of the muscle strip is warmed that end may recover first regardless of the original point of excitation. Cooling slows recovery. By cooling one end of the muscle strip one can delay recovery at that region. By decreasing the oxygen supply to a region of muscle it becomes hypoxic and recovery in that region will be delayed.

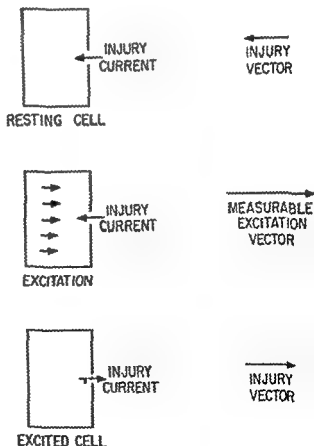


Figure 26

Drugs can alter recovery by their chemical action on the cell membrane. Changes in the electrolyte content of the cellular media and in the fluids surrounding the media will alter the usual driving forces across the cell membrane and may profoundly affect both recovery and excitation.

EFFECTS OF CELL INJURY

Cell injury creates a constant force at the area of injury. Injury connotes the destruction of the cell membrane. This permits a constant diffusion of electrical current between the internal and external environment of the cell. In the resting state the external environment is relatively positive; therefore a constant flow from outside to inside exists through the area of in-

jury (Fig. 26). Moreover the force across the membrane (resting membrane action potential) opposite the area of injury is unopposed. These forces create a vector towards the normal tissue area and away from the area of injury. The excited cell has a positive internal environment and creates a force toward the exterior. The membrane action potential is also reversed in the excited state. The resultant of these forces creates a vector toward the injured area during the excitation phase. Recording techniques are such that injury forces are measured only during the excited phase (between excitation and recovery). For this reason the injury vector is manifested only during excitation and the following rule may be formulated: *Injury creates a vector directed toward the area of injury.*

III

The Heart is a Source of Electrical Forces

ATRIAL EXCITATION

THE ATRIA are thin walled structures. The right atrium forms the right border of the heart and is anterior to the left atrium. The left atrium is the most posterior part of the heart and forms the right posterior chamber. Both are composed of muscle fibers giving them wall thickness. Like all resting cells a positive electrical field comprises the external environment. Of necessity both atrial surfaces (internal and external) are positive with respect to the cellular interior during the resting state.

Excitation of the atria begins from a stimulus emitted by the sino auricular node. This is a body of specialized conduction tissue situated at the orifices of the vena cava. It is a long narrow body. It is normally responsible for cardiac rate and is called the pacemaker. The node is subject to influences from the vagus nerve (chiefly the right vagus) and the sympathetic nerve supply. The former slows the node's activity and can arrest its action completely while the latter accelerates its activity.

The SA node is external to the atria causing excitation to begin at the external surface. The wave of excitation spreads away from its point of origin. Its pathway over the external surface can be roughly described as a series of expanding concentric circles. Since the atria are thin walled structures only a short time is required for excitation to reach the internal surface under normal circumstances. Nevertheless the fact that time for transmission is required should not be overlooked. The time factor is of importance in pathological states. The early arrival of the wave of excitation at the inner surface prevents

the wave of excitation from obtaining any large area (Fig 27). This in turn prevents normal atrial excitation from creating a large force. Should transmission to the inner surface be delayed the wave of excitation increases in size creating a larger force. Those clinical conditions causing delay in transmission of the wave front are associated with increased electrical force regardless of the presence or absence of atrial hypertrophy.

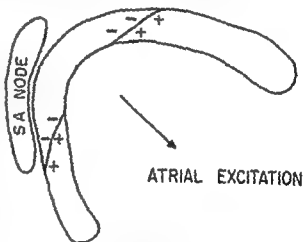


Figure 27

The force created by atrial excitation begins with zero magnitude, increases to its maximum and returns to zero. The resultant vector rotates its direction. The vector terminus of necessity describes a pathway in space or a loop. This loop represents atrial excitation and is called the P loop. Atrial excitation involves the successive addition and subtraction of vector quantities. The spatial pathway then becomes an expression of magnitude of the forces of excitation. The pathway is small as forces are being added

and subtracted simultaneously. This is the result of the arrival of the excitation wave at the inner surface prior to completion of excitation of the external surface.

Since atrial excitation spreads away from the SA node its force is directed away from the orifices of the vena cava. This is the old rule that the force of excitation is directed away from the excited area.

The time required for atrial excitation does not exceed 12 sec. The time interval depends upon the speed of radial transmission of excitation through the muscle fibers (3.5 mm per 0.1 sec) and the surface area of the atria. When the atrial volume is large the surface area to be excited is greater thus prolonging atrial excitation.

VENTRICULAR EXCITATION

In order to appreciate the spatial direction of forces created by ventricular excitation certain simple anatomic facts must be considered. 1) The left ventricle comprises the larger part of ventricular mass. It is shaped much like a cone. The base of this cone faces the right posterior chest. The apex of the cone is directed anteriorly and to the left. The cone rests on its side above the diaphragm. The wall of the cone is thickest at its base and becomes progressively thinner towards the apex. 2) The wall of the left ventricle is normally three or four times as thick as the wall of the right ventricle. 3) The margins of the free wall of the right ventricle are attached to the anterior surface of the left ventricular cone. The portion of the left ventricular wall bounded by this attachment represents the septum or partition between the ventricular cavities. 4) The septum is anatomically and functionally part of the left ventricle. The segment of the cone comprising the septum is a 60° arc. 5) The septum faces the anterior chest wall. The right ventricle then is anterior to the left ventricle (Fig. 28).

At the base of the septum next to the atria is the AV node. This tissue is a bundle of specialized conduction tissue which transmits the electrical impulse from the atria to the ventricles. Normally there is a delay between atrial and ventricular excitation due to the time required for the transmission of the excitation impulse through this conduction pathway to its receptor end plates in the myocardium. The node has many ramifications of tissue fibers spreading over both surfaces of the septum and the entire endocardial cavities of both ventricles. This tissue transmits impulses very rapidly. It permits rapid excitation throughout the endocardial surfaces.

The order of excitation of the component portions of the ventricle is now relatively well understood. 1) The left endocardial surface of the septum is activated first. 2) About 0.05 sec after the onset of excitation the right endocardial septal surface is activated. 3) The remainder of both endocardial surfaces complete activation by 0.2 sec after the onset of ventricular excitation. Thus at 0.2 sec excitation appears as two confluent cones of activity placed side by side. 4) Excitation spreads externally through the muscle thickness at a rate of 1/3 meter per sec (3 mm per 0.1 sec). 5) The thin walled right ventricle completes activation first normally between 0.20 and 0.40 sec. 6) The septum is activated from both right and left surfaces completing activation in less than one half the time required for complete ventricular excitation. For an excitation cycle of 0.08 sec duration septal activation should be complete by 0.4 sec. The septum creates a small force

The concept of rapid endocardial excitation is not new. It was originally proposed by Einthoven (1908). Wilson made use of Einthoven's explanation and his comments relative to endocardial and septal activation are as follows:

Excitation of the subendocardial muscle begins at many different points almost simultaneously (6) so that many islands of active tissue one for each junction between the Purkinje tissue and the ordinary muscle are quickly formed. As these islands grow they coalesce to form larger islands until both ventricular cavities are lined almost everywhere except at the orifices by a sheet of active muscle. At this stage of ventricular excitation there is a boundary between resting and active muscle in almost every part of the outside

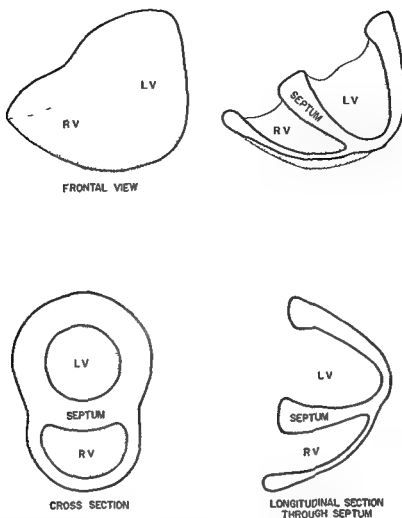


Figure 28

of very short duration because it is activated nearly simultaneously from both sides. After completion of right ventricular and septal activation, the wave of excitation is only part of a cone of activity located in the left ventricle. It is a cone with an arc of 60° cut away from its wall. The 60° arc represents the septal region that has completed activation. 7) The cone

or wave front becomes progressively shorter as new muscle areas complete activation. The cone is thinnest at the apex; thus the wave front arrives first at the surface of the left ventricle near the apex. It arrives at the surface at progressively later intervals from the apex to the base. The size of the wave front then is diminished from the apex to the base (Fig. 29).

The effective force of a wave of excitation is the resultant of the doublet vectors acting perpendicular to the wave front. With this principle in mind, the resultant vector of the wave of excitation can be determined at any stage of the cycle. 1) The left septal surface begins activation, creating a force directed away from the excited region. Considering the anatomic

walls in the septum which is activated from both sides there are two such boundaries. The two groups of electrical forces associated with the two septal boundaries are opposite in direction and cancel one another more or less completely so that the voltage across the septum is normally small. Wilson F. N., Rosenbaum F. F., and Johnston F. D. Interpretation of the Ventricular Complex of the Electrocardiogram. *Advances in Internal Medicine*. New York: Interscience Publishers Inc. 1947. Vol. 2, pp. 1-63.

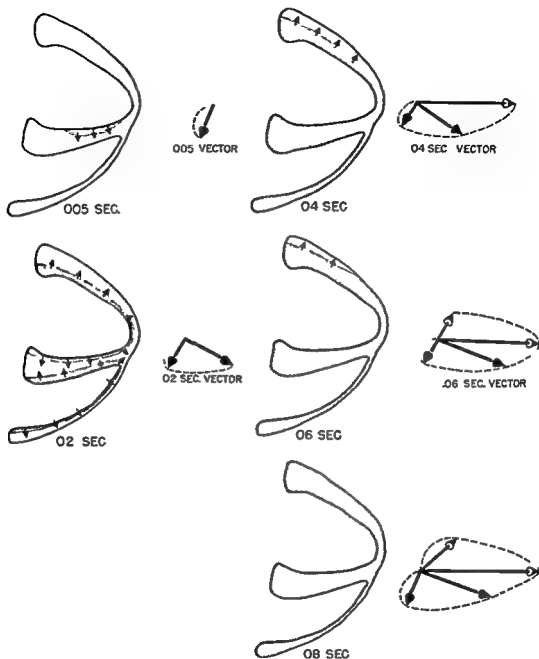


Figure 29

location of the septum this means the initial force is anterior and to the right. The duration of this unopposed force is 005 sec. At this time activation of the right septal surface begins and neutralizes the force from the left surface. 2) Endocardial spread continues in both cavities. When both cones are complete (02 sec) the flat surface area is bounded by the cones of excitation which is roughly equivalent to the

base of the ventricles. The resultant vector is proportional to this area and perpendicular to the base of the heart. It should be parallel to the long axis of the ventricles directed to the left and anteriorly. 3) Completion of activation of the right ventricular wall (035 sec) causes posterior rotation of the resultant vector. 4) As the wave front diminishes in size from apex to base the resultant vector magnitude decreases.

The vector rotates backward as it becomes shorter. Its terminal location is roughly perpendicular to the long axis of the left ventricle.

The instantaneous vector at the completion of right ventricular activation is of special interest (usually about 0.35 sec and usually the longest instantaneous spatial vector). It is the resultant of vectors representing two flat surface areas: the area of the base of the left ventricle and the flat surface area of the septum (Fig. 30). The term *long left ventricular vector* will be used to designate this vector. When the ventricle is a short cone the septal surface area is small and the long left ventricular vector is more nearly parallel to the long axis of the ven-

tricle further explored in the discussion of volume conductors.

The rotation of a central resultant on its point of origin is caused by the successive addition or subtraction of new vectors. The spatial pathway is a measure of the magnitude of newly added or subtracted forces. The pathway is the sides of a polygon. Because the events are rapid changes of small magnitude it becomes a loop. The magnitude of the force causing rotation from one central resultant position to another is the length of the spatial pathway. These principles are readily applied to ventricular excitation. The magnitude of the force acting to rotate the resultant vector from one position to

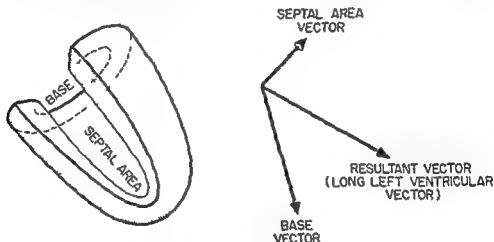


Figure 30

tricles. As the ventricle elongates the septal area is increased causing the left ventricular vector to rotate more posteriorly away from the long axis of the ventricle.

The continuous rotation of the resultant vector describes a spatial pathway. The resultants are thought of as rotating upon a center of origin. For ventricular excitation this center is confined to a small area in the left ventricle. Its stationary characteristic is due to the shape of the wave front of excitation. For the most part the wave front represents a cone or a 300 arc of a cone. With all forces acting perpendicular to the wave front it is easy to understand why the lines of force all intersect a relatively small area. The validity of this concept will be

another is equivalent to the distance between the two vector points or that segment of the spatial pathway. The length of the pathway between the terminus of the 0.4 sec vector and the 0.5 sec vector represents the change of force acting between the 0.4 sec interval and the 0.5 sec interval. This force is proportional to the change in the area of the wave front between the 0.4 and 0.5 sec intervals.

The length of the spatial pathway between the long left ventricular vector and completion of ventricular excitation is an index of left ventricular surface area (and volume). This measurement can be determined by indirect means and will be called the *index of maximum potential for the left ventricle*. It has a direct

relationship to the long left ventricular vector

The apical segments of the wave of excitation have a shorter duration of action than the basilar region. The duration of action for the conical wave front segment lost from 04 to 05 sec averages 04 to 05*. To express the potential in terms of duration of action represented by this conical segment in time units one can multiply the length of the spatial pathway from 04 to 05 sec by 045 sec. The product is expressed in $mv \times sec/100$.

An expression of the *potential seconds* for the conical segment represented by the time interval 05 to 06 can be obtained in a similar manner as above. After determining the $mv \times sec/100$ for each conical segment they may be added. The sum is an index of the potential $\times sec/100$ created by left ventricular excitation and may be called the *potential seconds for the left ventricle per cardiac cycle*. It is an expression of the surface area of the ventricle the thickness of the muscle wall and the density of electrical charges across the wave front†.

There is a marked increase in wall thickness at the base of the ventricles. This causes slow changes in the size of the wave front and terminal slowing of the spatial pathway described by the rotating instantaneous spatial resultant vector. This area creates further complications because it is not always well supplied with specialized conduction tissue. As a result activation may be somewhat delayed and create terminal variations in the direction of excitation forces.

VENTRICULAR RECOVERY

The isolated muscle strip begins recovery at the same site as the onset of excitation. The ventricle behaves in an exactly similar fashion. The thin walled right ventricle begins recovery

Actually 01 or 015 sec should be subtracted from the time interval for the time required for endocardial activation but for practical purposes this small period of delay may be ignored.

†Actual calculation of total potential will require conversion by geometric factors related to the length of the ventricle and the area of the base as used in conic calculations.

at its apex shortly or immediately after completion of ventricular excitation. The apex region of the left ventricle then begins recovery at its endocardial region*. Recovery proceeds from apex to base. The usual time required for recovery is between 26 and 36 sec depending upon cardiac rate.

Since the force across a wave front is always directed towards the resting (recovered) tissue the spatial forces of ventricular recovery can be analyzed. Initial apical recovery of the right ventricular wall creates a force anteriorly and leftward. This force is small because the area of the wave front is small. Recovery of the left ventricle from apex to base creates a force directed away from the base to the left and anterior. The magnitude of the force depends upon both the shape and size of the wave front of recovery. Normally it is smaller than the forces of excitation. The forces of recovery are roughly parallel to the long axis of the heart.

The important point about ventricular recovery is its onset at the apex. The base apex relationship to ventricular recovery was clearly demonstrated by Mines† in 1913 by means of a base apex lead. He observed that warming of the base of the heart created a negative T wave. A negative T wave was made positive by apical warming. Cooling of the apex produced a negative T wave. He correctly attributed these findings to change in speed of recovery, increased with warming and slowed with cooling. Later Wilson and Herrmann (1921)§ demonstrated

The initial recovery of the endocardium is responsible for the negative force of recovery following the upright excitation wave as recorded directly from the surface of the heart muscle. Sir Thomas Lewis demonstrated this pattern and called it an electrogram. Recent interest in this observation has been revived since its frequent demonstration during cardiac surgery. The area of muscle beneath the electrode is to be regarded as an isolated muscle strip. Lewis Thomas *The Mechanism and Graphic Registration of the Heart Beat*. New York: Paul B. Hoeber, Inc. 1920.

†Mines C. R. On functional analysis by the action of electrolytes. *J. Physiol.* 46:189, 1913.

§These classic experiments were done in experimental bundle branch block. The end of the refractory period was correlated with the decline in excitation. In right bundle branch block the left ventricle passed out of the refractory state 02 to 04 sec before the right ventricle. The deeper

that recovery followed the same general order as excitation

A popular theoretical explanation for the direction of the forces of ventricular recovery is delay in recovery of the endocardial regions. This concept is without experimental support. It is contrary to known experimental observations. As long as apical recovery occurs first the resultant vector will be directed away from the base regardless of whether recovery begins at the endocardial or epicardial surface (Fig 31)

Layers of the left apex passed out of the refractory state before the surface layers. The right ventricle recovered before the left ventricle in the presence of right bundle branch block. On the basis of these observations the authors stated:

We have shown that the order of recovery of the various regions of the ventricular mass is the same as the order of excitation. Wilson F N and Herrmann G R. An experimental study of incomplete bundle branch block and of the refractory period of the heart of the dog. *Heart* 8:239 1921.

Dr. Herrmann has been kind enough to review the concept of the base apex relation in production of the T wave and is in agreement with the concept set forth.

The concept of the ventricular gradient was ushered into electrocardiography as a pure theoretical paper presented by Wilson in 1931. It is of interest to note that he defined the ventricular gradient as the mean electrical axis of Q R S T which pointed away from the ventricular region with the longest average duration of excitation and towards the region of least average duration. He pointed out that the normal ventricular gradient pointed in a base apex direction because the average length of systole was greater at the base than at the apex. Wilson F N, MacLeod A G and Barker P S. The T deflection of the electrocardiogram. *Tr A Am Physcians* 46:29 1931.

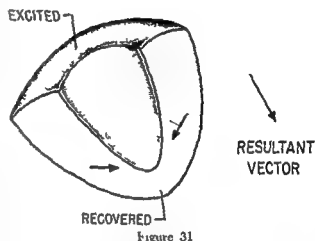
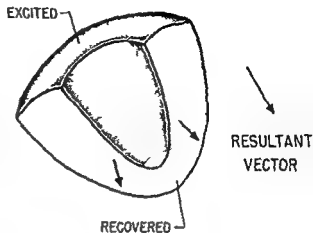


Figure 31

The wave front of recovery closely resembles a complete conical segment (360°). A change in the inclination of the walls of the cone greatly alters the resultant vector magnitude. A cylin-

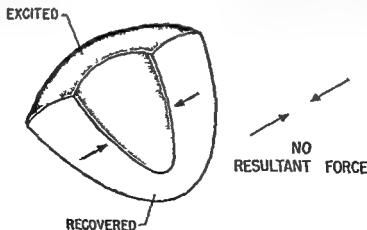


Figure 32

dical wave front creates forces opposed to each other. The resultant approaches zero magnitude. This is one explanation for demonstration of excitation forces and absence of demonstrable recovery forces (Fig. 32). The creation of an everted conical wave front reverses the customary direction of recovery forces. The resultant vector is directed toward the base rather than toward the apex (Fig. 33).

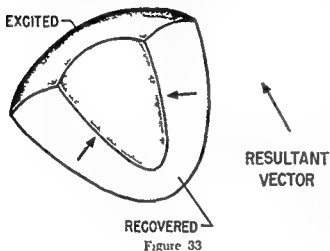


Figure 33

The normal course of recovery creates a continuously changing vector describing a loop. The loop is enclosed within the QRS loop and is smaller in magnitude. Commonly it is somewhat anterior to the point of origin. It is called the T loop. Its spatial pathway is a function of the magnitude of the component forces. Since there is a continuous loss of forces at the same time of addition of new forces the spatial pathway is small in comparison to the QRS pathway. The T loop appears somewhat anterior to the QRS loop because the latter has a long posterior course caused by early completion of septal activation.

AFTER POTENTIAL

A small additional wave of electrical activity may follow ventricular recovery. This event is called the U event. Its cause is not completely understood. It will also describe a spatial pathway and its forces are roughly parallel to the forces of ventricular recovery.

IV

Fundamentals of Conductors

CHARACTERISTICS OF CONDUCTORS

A CONDUCTOR is a medium for the transmission of electricity. When a complete circuit exists current flows through a conductor. The media serving as a conductor offers resistance to the transmission of electrical force. When the resistance is great the forces are poorly transmitted. Such a media is a poor conductor. A media offering little resistance to transmission of forces is a good conductor. In a general sense all substances are conductors. However some are such poor conductors that a very large force must be present to transmit electrical forces any great distance from the source. Such substances are called insulators. Wood and rubber are examples of poor conductors. Copper and zinc are examples of good conductors.

Conductors may be either solid gas or liquid. A volume of liquid used as a conductor is called a volume conductor. Water is a good conductor when it contains electrolytes the greater the concentration of electrolytes the better its conductivity.

When a conductor offers a uniform amount of resistance to electrical forces throughout it is called a *homogeneous conductor*. In such a conductor if one knows the amount of resistance offered by 1 centimeter of distance from the source one can calculate the total resistance offered at any distance from the source. A conductor which does not offer a uniform resistance throughout is called a *heterogeneous conductor*. Due to the variation in resistance at different areas the total resistance cannot be determined in the same manner as for a homogeneous conductor.

ELECTRICAL FIELD WITHIN A VOLUME CONDUCTOR

An electrical field is created around a charged particle within a volume conductor. Potential diminishes as the distance from the particle (source) is increased. In volume conductors the potential diminishes in proportion to the square of the radius distance (r). This is due to the spherical dissipation of the force. In a flat disk conductor potential diminishes simply in relation to the radius distance (r).

The presence of two fields of unequal potential (positive and negative) creates a resultant force. The direction of this force is a straight line drawn from the point of lowest potential (negative pole) to the point of greatest potential (positive pole). The force may be represented as a vector directed toward the positive pole and away from the negative pole. This vector represents the axis of the electrical field. It is the dipole vector (or doublet vector). The magnitude of the dipole vector depends upon the difference in potential between the two poles and is called the dipole moment (electrical moment).

In electrocardiography we are concerned with a negative field on one side of a wave front and a positive field on the opposite side. The magnitude of the dipole moment (electrical moment) depends upon the density of charges (doublet vectors) and the flat surface area of the wave front. It is then an expression of the effective force created by the wave front.

INFINITE VOLUME CONDUCTORS

At any point within an infinite volume conductor the potential depends upon 1) the mag

magnitude of the dipole vector or electrical moment M 2) the distance of the point from the center of origin r and 3) the angle between the dipole vector and the coordinate drawn from the potential point and the center of origin (Fig 34). This may be expressed as a simple equation

$$\text{Potential} = \frac{M \cos \theta}{r}$$

If one measures the potential and knows angle θ and the radius distance the dipole moment M can be determined

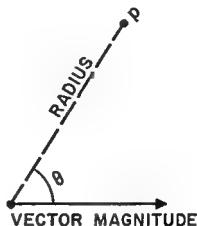


Figure 34

FINITE VOLUME CONDUCTORS

The potential is always greater at the surface of a limited volume conductor than it would be at the same distance from the source in an infinite volume conductor. This is due to the insulating effect of the surface. These effects are not so noticeable near the center of a tank. For a spherical type tank it has been determined that the potential is approximately three times as great at the surface as would be expected for that distance. This is particularly important in electrocardiography as measurements are made from the surface. The potential at the surface of a spherical finite volume conductor is expressed

$$\text{Potential} = 3 \times \frac{M \cos \theta}{r}$$

The shape of the boundary also influences the potential measurement. A rectangular tank will distort the apparent direction of the dipole vector. Variations in resistance within the volume conductor will further distort the direction of the lines of force and alter potential measurements.

Wilson F N, MacLeod A G and Barker P H. *The Distribution of the Currents of Action and Injury Displayed by Heart Muscle and other Excitable Tissues*. Univ of Mich Studies Scientific Series Ann Arbor Univ of Mich Press 1933 Vol 18 p 58.

"at any point upon the surface of the sphere the magnitude of V is three times as great as it would be at the same point in an infinite medium."

THE BODY AS A VOLUME CONDUCTOR

For practical application it has been found convenient to consider the body as a homogeneous volume conductor. The body consisting primarily of water and electrolytes is a good volume conductor. It is true that resistance is not uniform throughout the body. Even the resistance offered by the skin varies at different points. The change in the character of the tissues alters the resistance factor. The air in the lungs is considered to act as an insulator. On the other hand the lungs are highly vascular and vascular beds are extremely good conductors. There are sources other than the heart creating electrical fields. In actuality then the body is a heterogeneous volume conductor of variable resistances with multiple sources of electrical fields. In application considering the body as a homogeneous volume conductor has been surprisingly satisfactory. Since this is a simple practical approach it shall be so considered here.

The wave front sets up an electrical field in the body. The isopotential plane defined on the body is perpendicular to the resultant vector of the wave front. The potential at any one point on the body is expressed by the equation

$$\text{Potential} = 3 \times \frac{M \cos \theta}{r^2}$$

By locating the isopotential plane one can esti

mate the direction of the resultant vector. It is perpendicular to this plane. Errors due to variation in resistance and shape of the finite boundary are assumed.

LOCATION OF THE ZERO CENTER

A perplexing problem in electrocardiography concerns the location of the zero center within the volume conductor (body). Much of the earlier work was based on the principle of the centric placed zero center. Ample evidence exists proving the zero center is actually eccentric in position. Moreover its eccentric location is a large factor in measuring potential at the surface of the body. There is no system of electrocardiography currently employed that avoids this error. This point will be discussed further in considering the lead systems.

Although considerable variation exists the zero center for ventricular excitation is usually three to four centimeters left of the mid sternal line about the lower level of the fourth intercostal space and one third the distance of the anterior posterior diameter of the chest from the sternum (Fig 35). The anterior location of the

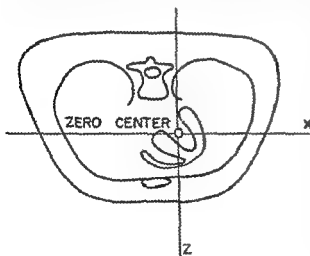


Figure 35

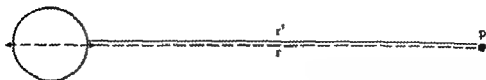


Figure 36

center should be no surprise to those who have examined a sagittal X-ray of the chest. The mediastinum is clearly seen to occupy the anterior one third compartment the remainder being composed of muscle and bone.

REMOTE ELECTRODE AND PARTIAL LEAD EFFECT

Another complication in measuring potential from the surface of the body arises from the so called "partial lead" effect. The implication is that potential values at points close to the heart are largely influenced by the nearest segment of muscle rather than by the entire muscle mass. A point sufficiently distant from the heart not to be influenced by proximal muscle segments is called remote. Both the diameter of the heart and distance from the heart determine whether a site is remote or not. Consider two doublet vectors of the heart in relation to a point P (Fig 36). When the diameter of the heart is small the difference in radius distance from P to the doublet vectors is proportionally not so great.

The effect of distance is enhanced by resistance. A conductor of low resistance would require greater distances from the source to be a remote site. Experiments on the isolated perfused mammalian heart immersed in a volume conductor have been carried out.* Such experiments indicate that beyond distances of two to three times the diameter of the heart the site can be considered as remote. These studies verify the concept of a center of origin for the forces created by the heart when recorded at distances great enough to be considered remote.

Hartmann I, Veyrat R, Weiss Oscar A M and Duchosal P W. Vectorcardiography as studied on the isolated mammalian heart suspended in a volume conductor. *Cardiologia* 129: 1935

Fundamentals of Electrocardiographic Instruments

THE PREVIOUS DISCUSSIONS have indicated that electrical fields are created by the heart as a source in a volume conductor the body. At all points on the body surface potential exists due to the cardiac electrical cycle. These areas of potential have values which have a direct relationship to the heart and the character of the volume conductors. When the potential values on the body surface are known the electrical activity of the heart can be ascertained. The next step is logically an instrument capable of measuring these potential values. To this end three major types of recording devices are employed the string gauge galvanometer with photographic recording system the galvanometer with direct writing recording system and the cathode ray oscilloscope.

STRING GAUGE GALVANOMETER

The string gauge galvanometer consists of a conductor suspended between the two poles of an electromagnet. The conductor is a quartz glass fiber coated with platinum. The electromagnet sets up an electrical field. When an electric current passes through the conductor the string is moved up or down in accordance with the principles of Fleming's law. One can illustrate this principle with the right hand (Fig 37). The third fourth and fifth fingers are pointed in the direction of force between the two poles of the electromagnet. The index finger is pointed in the direction of current flow through the suspended conductor. The extended thumb indicates the upward displacement of

the conductor. If the direction of current is reversed through the conductor it is deflected downward.

If electrical flow enters one terminal of the conductor a positive or upward displacement occurs. This terminal of the conductor is designated as the positive terminal or the positive

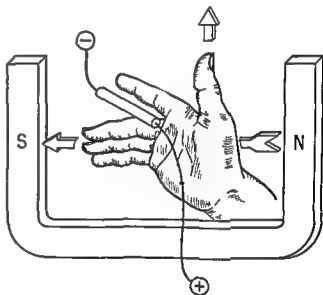


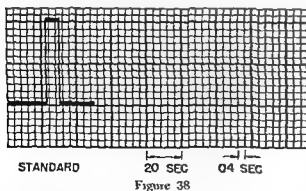
Figure 37

electrode. Current entering the other terminal of the conductor causes a downward displacement. This terminal is designated as the negative electrode. The amount of displacement of the filament is dependent upon the difference in potential at each end of the conductor. The principle of the galvanometer may be summarized as follows: A positive force entering the positive terminal (electrode) records a positive displacement.

The string gauge galvanometer is considerably more reliable than the direct writing instruments and for precise work it is much superior. It has been replaced largely because of the difficulties encountered in maintaining its day to day operation. The string properly adjusted offers little impedance to displacement and as a consequence there is little lag. To record the early events of septal activation lag should be less than 0.01 sec.

The string gauge galvanometer records deflections of the string upon a moving film strip. The film strip may be marked at time intervals by spoke wheels or prisms. Usually the film is moved at 25 mm per sec. Faster speeds are frequently an adjunct and 50 mm per sec is commonly utilized. A vertical marker is indicated on the film at each 0.4 sec interval. At each 20 sec interval a heavier marker is indicated. It is always the fifth vertical marker. At a speed of 25 mm per sec the markers are one mm apart. At 50 mm per sec they are two mm apart. In either case they represent 0.4 sec.

The deflection of the string is the means of measuring the magnitude of the force for any lead. For standardization one mm is applied to the conductor. At full standardization this produces one cm displacement of the string.



This is an index of the sensitivity of the galvanometer (Fig 38). When the sensitivity is known any displacement of the string can be measured in millivolts. If it is not known such

measurements cannot be made. Frequently portions of records are recorded at half standards one millivolt producing $\frac{1}{2}$ cm deflection.

THE DIRECT WRITING INSTRUMENT

The direct writing instruments utilize the principle of the string gauge galvanometer but the conductor and writing arm are heavy and the contact of the writing arm itself creates impedance to displacement. In a number of clinical conditions this is a serious handicap as it may obscure significant Q waves or small R waves. The peak of response to a force may be delayed from 0.1 to 0.15 sec. From the earlier discussion of rapid ventricular excitation it is clear that such measurements leave much to be desired. The direct writing instrument utilizes paper that is already graphed; the vertical lines are one mm apart. At a speed of 25 mm per sec each square equals 0.4 sec. The writing arm should not be permitted to record in the bottom or top $\frac{1}{2}$ cm of the graph for most instruments. The swing of the writing arm is damped at the limits of its excursion. The damping distorts (diminishes) the actual magnitude of the deflection.

CATHODE RAY OSCILLOSCOPE

The third type of recording instrument and in my opinion the best is the cathode ray oscilloscope (Fig 39). It is a good instrument and has an excellent frequency response and no appreciable lag. The cathode ray tube emits an electron beam to the screen of the tube. On one side of the beam is a positive plate and opposite it is a negative plate. Any change in potential on the plates causes a movement of the beam. A second pair of plates is placed perpendicular to the first pair. Either pair may be used to record the potential difference between two points. The instrument may be standardized (per mv) in a manner similar to the string gauge galvanometer. In addition to re-

The Cambridge Simple Cathode Ray Oscilloscope

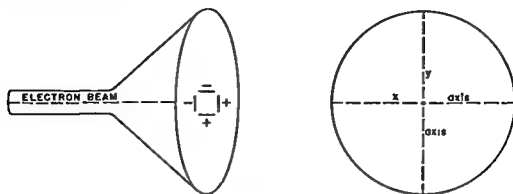


Figure 39

Recording an ordinary ECG the presence of two perpendicular pairs of plates permits the use of this instrument for recording vector cardiograms which will be discussed more fully at another point in the text

As a minimum standard any of the instruments should have the following specifications

1) Frequency response With an input volt

age of 15 cycles per sec an output response of 90% of the input should be obtained With an input voltage of 40 cycles per sec the output response should be 80% of the input

2) Linear response When one mv is applied the peak response should be reached at 0.1 sec It should be maintained at this peak plus or minus 10% for 0.2 sec

VI

Fundamentals of Electrocardiographic Leads

By UTILIZING the instruments for measuring potential at different points one can obtain an indirect measurement of the electrical events of the heart. The points selected for potential measurement have been standardized and constitute the lead systems used in electrocardiography. Any lead may be analyzed if the basic principles of potential measurement are properly applied.

EINTHOVEN'S LEADS

Einthoven originally measured potential at three points: left arm, right arm, and left leg. The difference in potential between these three points constitute Leads I, II, and III. It was assumed that all three potential points were equidistant from the zero center and equally distant from each other. Such leads are true bipolar or balanced leads. The extremities are considered as extensions of the lead wires. Thus the potential is measured at the surface of the trunk (the volume conductor). The potential (V) points are then designated as from each shoulder (V_L and V_R) and foot (V_F).

Given a spatial vector with a point of origin equally distant from the potential points V_L minus V_R will measure only the potential difference proportional to the X component of the spatial vector. The Y component of the spatial vector has the same effect upon positions V_L and V_R. If V_R is a negative electrode and V_L a positive electrode, they cancel each other according to no potential difference due to the vertical component: $-5 - (-5) = 0$. The same principle applies to the sagittal component. Such a lead measures only the X component of

the spatial vector and it is an X coordinate (Fig. 40). The magnitude of the vector measured by the lead is equal to X vector component magnitude times $\frac{1}{r}$, where L is the length of the lead (distance between V_L and V_R) and r

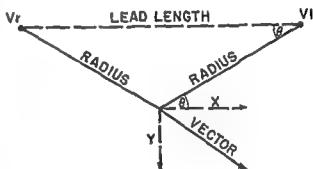


Figure 40

is the radius distance between the point of origin and V_R or V_L.

$$\text{Potential} = \text{Vector Magnitude} \times \frac{\cos \theta}{r}$$

$$\cos \theta = \frac{L}{r}$$

or the potential at a given point V_L equals

$$3 \left(\frac{L}{r} \times \frac{1}{r} \right) = 3 \left(\frac{L}{r^2} \right)$$

$$\text{the potential at } V_R = -3 \left(\frac{L}{r} \right)$$

$$V_L - V_R = 3 \left(\frac{L}{r} \right) - \left(-3 \times \frac{L}{r} \right) \text{ or } 3L/r$$

The derivation of these simple equations demonstrates that a true bipolar lead measures potential magnitude in accordance with the radius distance and lead length. When these measure

This is from the formula of point potential at the surface of a spherical volume conductor.

ments are known it is possible to determine the relative magnitude of the vector component as it exists at the point of origin within the volume conductor *

EINTHOVEN'S TRIANGLE

Considering all three potential points (Vr, V_f, V_i) is equally distant from the zero center and of equal distance from each other they describe an equilateral triangle (Fig 41). This is called Einthoven's triangle. Lead I with a positive electrode at V_i and a negative electrode at

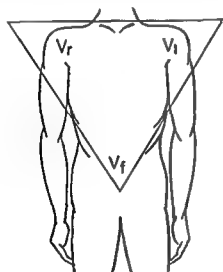


Figure 41

Vr measures potential V_iVr. Lead II with a positive electrode at V_f and a negative electrode at Vr measures potential V_fVr. Lead III with a positive electrode at V_f and a negative electrode at V_i measures potential V_fV_i. The Einthoven triangle describes a circumstance wherein 3Lr is the same for each lead. Each lead is a coordinate.

EINTHOVEN'S LAW

A relationship exists between the three leads which is expressed as Einthoven's law. Lead I + Lead III = Lead II. This may be stated (V_iVr + V_fV_i) = (V_fVr). Einthoven's Law may be proved by assigning arbitrary

values for the potential at each of the three points. If the potential at V_i is +5, Vr -10 and V_f +15, the law states:

$$\text{Lead I} = V_i V_r = +5 - (-10) = 5 + 10 \text{ or } +15$$

$$\text{Lead III} = V_f V_i = +15 - (+5) = 15 - 5 \text{ or } +10$$

$$\text{Lead II} = V_f V_r = +15 - (-10) = +15 + 10 \text{ or } +25$$

THE BIPOLAR TRIAXIAL REFERENCE SYSTEM

The leads comprising Einthoven's triangle may be rearranged geometrically to form a Triaxial Reference System. Each of the three leads are bipolar with a positive and negative electrode located at different points on the volume conductor. At a point between the two electrodes there is an isopotential point. This point is located by constructing a perpendicular from the lead axis to the center of zero potential. The perpendicular intersects the isopotential point or the point of origin for the lead. If the three points of origin for Leads I, II, and III are superimposed a Triaxial Reference System is formed (Fig 42). Since the original triangle

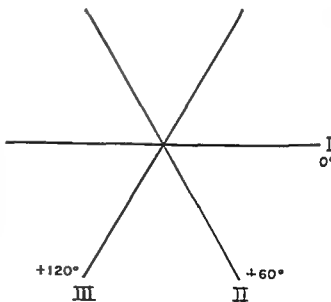


Figure 42

was equilateral with angles of 60°, the angles formed about the center of origin of the Triaxial Reference System will also be 60°. The point of origin of the Triaxial Reference System is considered as superimposed on the center of zero

* This is a relative value; resistance is not known.

potential within the chest. This is valid as each lead has the same relative sensitivity i.e. each lead records $3L/r^2$ times the actual force acting at the zero center.

For purposes of uniform orientation Lead I has been designated as an axis of zero degrees. All points below Lead I are spoken of as positive degrees away from Lead I. All points above Lead I are spoken of as negative degrees away from Lead I. Thus the positive pole of Lead II is $+60^\circ$ and the positive pole of Lead III is $+120^\circ$.

LAW OF PARALLELOGRAMS

The direction and magnitude of a vector originating from the zero center can be calculated from its projection on any two leads. This is done by utilizing the Law of Parallelograms. If the vector registers a positive value of $+5$ in Lead I and a value of $+2$ in Lead III, this is done as follows: 1) Mark off five units from the point of origin towards the positive pole of Lead I. 2) Construct a perpendicular from Lead I at this point. 3) Mark off two units from the point of origin towards the positive pole of Lead III. 4) Construct a perpendicular from Lead III at this point. The two perpendicular lines will intersect one another. 5) A line is drawn from the point of origin of the reference system to the point of intersection by the two perpendicular lines. This line represents the direction and magnitude of the unknown vector. The vector is the diagonal of a parallelogram (Fig. 43).

The above process may be reversed. If the direction and magnitude of a vector is known one can calculate its projection on any one of the three lead axes. The direction of the vector is defined in degrees away from the axis of Lead I.

A vector with a magnitude of $+6$ and an axis of $+60^\circ$ will be parallel to Lead II. It may be represented by marking off 6 units upon Lead II. A line constructed from this point on Lead II to a point perpendicular to Lead I will designate 3 positive units on Lead I. A second

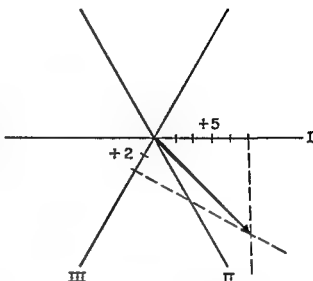


Figure 43

line drawn from the terminal point of the force on Lead II to a point perpendicular to Lead III will designate 3 positive units on Lead III. The value of the vector projected on both Lead I and Lead III will be $+3$ units, conforming to Einthoven's Law $I + III = II$ or $+3 + 3 = +6$. This also illustrates an important characteristic of a vector: *the lead most nearly parallel to a vector will record the largest deflection* (Fig. 44).

A vector with an axis of $+90^\circ$ and a magnitude of 6 units will record 5 units in Lead II.

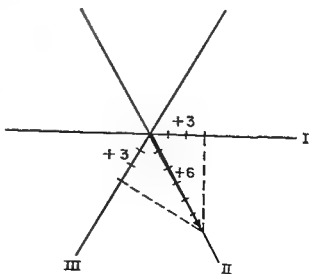


Figure 44

and 5 units in Lead III. The perpendicular from Lead I to the tip of the vector will fall at zero on Lead I. Such a vector will record no potential in Lead I. This satisfies Einthoven's Law as well as the Law of Parallelograms. Another important vector principle is illustrated. When the axis of a vector is 90° away from the lead axis, no potential is measured by that lead. Such a lead is called a diphasic or isoelectric lead (Fig. 45).

A vector with an axis of $+30^\circ$ (perpendicular to Lead III) records no potential in Lead III. A vector with an axis of -30° (perpendicular to

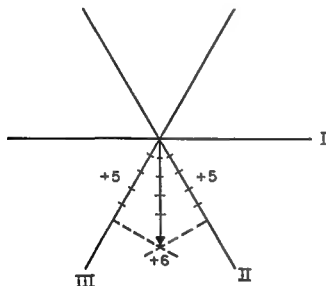


Figure 45

Lead II) records no potential in Lead II. The axis of a vector can be determined by inspection alone whenever one of the three leads is diphasic or isoelectric.

THE V LEAD

A bipolar lead measures the potential difference between two points. Utilizing the three standard leads, it is impossible to record the potential received at any one point due to the influence of the other electrode. Wilson devised a method of eliminating the mathematical value of the negative electrode. This was accomplished by attaching the negative terminal of the conductor to the right arm, left arm, and

left leg. The value received at the negative pole would theoretically be zero (Fig. 46).

The principle of zero potential at the negative electrode is illustrated as follows. Assume a vector with a magnitude of $+10$ and an axis of $+90^\circ$. The potential received by a negative electrode V_f would be -10 by the negative

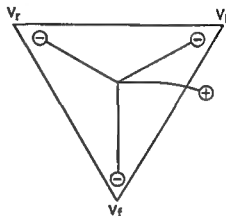


Figure 46

electrode $V_r -5$ by the negative electrode $V_f -5$. By algebraic addition $-(+10) - (-5) - (-5) = -10 + 5 + 5 = 0$. The positive electrode could then be placed at any point and measure only the potential received at that point. Such a lead is called a V lead or an exploring electrode.

THE UNIPOLAR TRIAXIAL REFERENCE SYSTEM

Placing an exploring electrode (V lead) on the right arm, left arm, and left leg creates leads V_r , V_1 , and V_f . The axis of Lead V_r may be depicted by drawing a line from the right shoulder to the center of zero potential. Lead V_1 has an axis from the left shoulder to the zero center. Lead V_f has an axis from the pubis to the center of zero potential. V_r , V_1 , and V_f bisect each angle of Einthoven's triangle. The negative terminal is the same for each lead and equivalent to zero. This point may be considered as the point of origin for the three leads. If the axis of each lead is extended beyond its point of origin, another Triaxial Reference System is formed with 60° between each axis (Fig. 47).

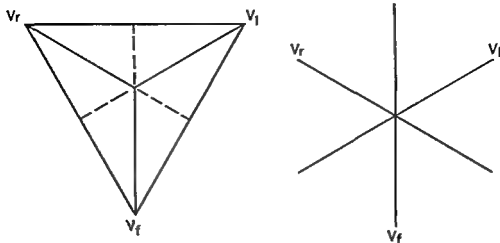


Figure 47

The unipolar limb leads also bear a mathematical relation to one another. This may be expressed $V_r + V_1 + V_f = 0$. If a force with an axis of $+90^\circ$ and a magnitude of 6 units is measured by all three leads, Lead $V_f = +6$, $V_r = -3$, $V_1 = -3$. This may be expressed $+6 - 3 - 3 = 0$.

HEXAXIAL REFERENCE SYSTEM

The Unipolar Triaxial Reference System is rotated 30° away from the Bipolar Triaxial Reference System. By superimposing the point of origin for both Triaxial Reference Systems, a Hexaxial Reference System is formed (Fig. 48). The positive electrode of V_r has an axis of -150° , V_1 -30° and V_f $+90^\circ$.

AUGMENTED UNIPOLAR LEADS

Because the influence of the negative pole is removed, the amplitude of the unipolar leads is only half that obtained by the bipolar limb leads. Goldberger amplified (augmented) the potential by removing the negative electrode from V_1 when the positive electrode was on the left arm and in a like manner on the other extremities. To indicate this augmentation, the prefix *a* was added before the symbol of each lead (aV_r , aV_1 , aV_f). Even with augmentation, the relative deflection obtained in the augmented limb leads is equivalent to only 86% of the value measured by the bipolar leads.

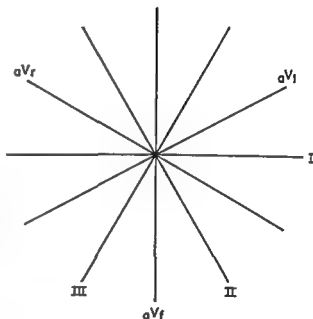


Figure 48

AXIS BY INSPECTION

If the unipolar limb leads were equivalent to the bipolar limb leads, the lead with the most positive deflection would be most nearly parallel to the vector axis. Regardless of the relative sensitivity of each lead, no potential will be registered at a point perpendicular to the vector (Fig. 49). This principle enables one to tell by simple inspection of the six limb leads the axis of any force. If no potential is recorded in Lead

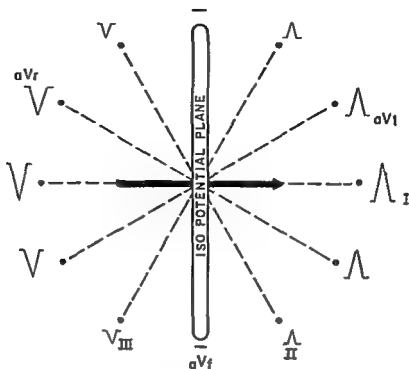


Figure 49

aVL (axis -30°) the axis of the vector is $+60^\circ$ (90° away from aVL). In this event Lead II will record a strongly positive force. The vector axis could have been -120° . In the latter case the force would be directed away from the positive terminal of Lead II and the complex in Lead II would be negative (Fig 50).

To summarize the above discussion the following rules may be formulated:

- 1) When a vector force faces a positive electrode a positive wave is recorded.
- 2) When a vector force is going away from

the positive electrode (towards the negative electrode) a negative wave is recorded.

3) The lead axis most closely parallel to the axis of a vector will record the greatest negative or greatest positive deflection (allowing for differences in value of the augmented unipolar and bipolar leads).

4) Ninety degrees away from a vector the mathematical value of the potential measured equals zero (this perpendicular plane is called an isopotential plane).

5) Utilizing the Hexaxial Reference System

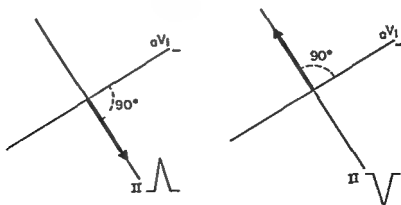


Figure 50

one may determine the axis of any vector by the following steps

a) Select the lead recording an isoelectric (mathematical zero) complex. The axis of this lead is the axis of the isopotential plane for the vector.

b) Ninety degrees away from the isopotential plane is the direction of the vector. If the lead that is parallel with the direction of the vector records a negative wave the axis of the vector is 180° away from the positive terminal of that lead.

When no isoelectric complex is recorded in

using this method the axis of a vector can be determined within 5° to 10° of its true axis.

A vector with an axis between +30° and +60° will record positive values in all leads except Lead aVr. The axis of the vector can be deduced. A vector of +30° records an isoelectric value in Lead III therefore the vector must have an axis of more than +30°. A vector of +60° records an isoelectric value in Lead aVI therefore the axis must be less than +60°. Whenever all leads except aVr record a positive value the vector must be between +30° and +60° (Fig. 52).

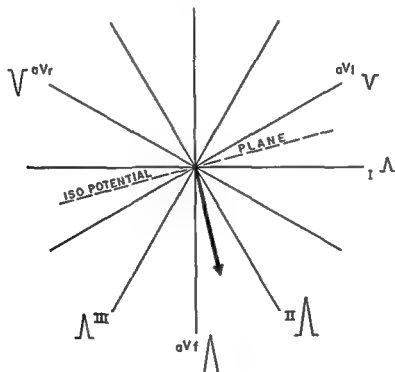


Figure 51

any lead the isopotential plane of the force can still be determined. This is done by noting the point on the Hexaxial Reference System where the recorded complexes change from positive to negative. If aVL records a negative complex and Lead I records a positive value the isopotential plane is between their lead axes (between -30° and 0°). One can arbitrarily locate the isopotential plane at -15°. Ninety degrees away from -15° would be +75° (Fig. 51). By

THE FRONTAL PLANE

The Hexaxial Reference System describes a plane around the body. This plane is flat like a sheet of paper and has no depth. If a vector is acting only in this plane its total value can be measured by the Hexaxial Reference System. The vector forces originating from the heart are also directed either anteriorly or posteriorly. The six limb leads measure only the projection of the spatial vector on the frontal plane. The

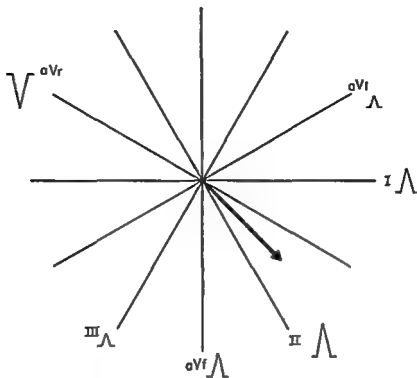


Figure 52

more anterior or posterior the vector the less will be its magnitude of projection on the frontal plane. The two perpendicular coordinates (X and Y) of the frontal plane are lead axis I and lead axis aVF .

THE TRANSVERSE PLANE

The projection of a cardiac vector on the transverse plane is measured by the V leads. This plane may be described as a cross section through the chest. The transverse plane consists of two perpendicular coordinates. One coordinate is parallel with Lead I and is the X coordinate. The other coordinate is the Z coordinate, measuring the anterior-posterior component of the vector.

The V lead electrodes used to measure precordial potential are as described previously for an exploring electrode. The location for each V lead comprising the transverse plane is at a specific anatomic site. The usual leads taken are at six sites around a little more than a quarter of the circumference of the chest. Their locations are as follows (Fig 53).

- V1 Fourth interspace to the right of the sternum
- V2 Fourth interspace to the left of the sternum
- V3 Halfway between V2 and V4
- V4 Fifth interspace left midclavicular line

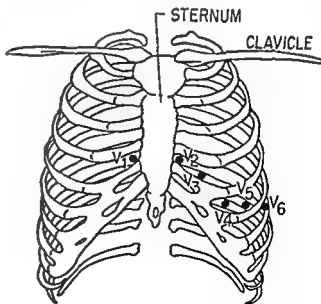


Figure 53

V5 The same horizontal plane as V4 and in the anterior axillary line

V6 The same horizontal plane as V4 and V5 in the mid axillary line

Occasionally it is desirable to measure potential at additional areas. Right precordial leads may be used. These leads are called VR leads. They are in exactly similar locations as the usual left chest leads only on the right side. V1R is the same as V2 position. V2R is the same as V1. V3R is comparable to V3 position only on the right chest rather than the left (Fig. 54).

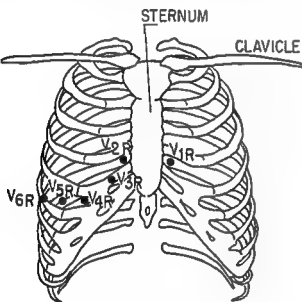


Figure 54

When the posterior axillary line position is utilized it is called V7. A point at the level of V6 and in the left tip of the scapula is utilized for V8. Just to the left of the spine on the same level is position V9.

The anterior precordial leads are not usually considered "remote." Their location is not sufficiently distant from the heart for a remote electrode. Their proximity results in discrepancies in amplitude. Nevertheless when the precordial electrode is located in the positive electrical field, positive force will be recorded and when in the negative electrical field a negative wave is recorded. This enables one to locate the isopotential plane for a cardiac vector.

The vector force between the two electrical fields is perpendicular to the isopotential plane. Locating the vector in the transverse plane is possible by first locating the isopotential plane on the chest.

When the V lead position is sufficiently distant from the heart to be considered "remote," the potential at its location is expressed simply as $\frac{3}{r}$. It differs from the previously described bipolar leads in that no angle exists between the lead axis and the radius distance as they are one and the same.

The proximity of the precordial leads to the zero center exaggerates small variations in the location of the center. While it is recognized that the zero center has individual variations its location can be approximated. Most authorities agree that it lies anteriorly to the anatomic center of the thorax. It is convenient and practical to consider its location at a point between V6 and V6R at the level of the fourth intercostal space and just to the left of the sternum. The zero center can then be located as the point where Leads I, aVF and V2 intersect and are mutually perpendicular. These three leads form three mutually perpendicular coordinates (X, Y, Z).

All V leads have the same theoretic negative pole, the zero center. They comprise an axis system for the transverse plane. The angle between each V lead axis will have individual variations but an approximation can be utilized.

Because the zero center is displaced anteriorly, it is considered on a plane between V6 and V6R positions (a plane between the mid axillary lines). Often V6 may be just a little posterior to the true plane of zero potential, however V6 position is more nearly correct than V5 which is usually anterior to the zero center plane.

V2 is one of the perpendicular coordinates and 90° away from V6. Lead axis V3 forms an angle of 20° with lead axis V2. Lead axis V4 forms a 40° angle with lead axis V2. Lead axis V5 forms a 70° angle with V2. The axis of Lead I is parallel to the axis of Lead V6. As a reference system in the transverse plane it is con-

venient to call V6 zero degrees. Every point posterior to V6 is considered positive. The transverse plane axis can be located as follows (Fig 55)

V1 = -110
 V2 = -90
 V3 = -70
 V4 = -50
 V5 = -20
 V6 = 0

THE TWELVE LEAD SPATIAL REFERENCE SYSTEM

If one superimposed the point of origin for both the Hexaxial Reference System and the Transverse Reference System a Spatial Reference System is formed (Fig 57)

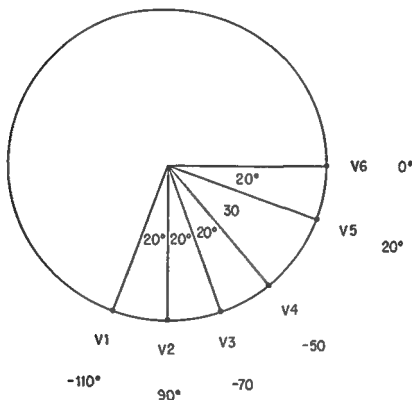


Figure 55

The magnitude of the vector projected on the transverse plane can be determined by the following steps (Fig 56)

- 1) Locate the isopotential point e.g. V3 at -70
- 2) The transverse vector is projected 90° away from that point i.e. +20° (Twenty degrees is the angle between the vector and the X coordinate or Leads I and V6)
- 3) Measure the mean value of the vector on its X axis in the frontal plane (Lead I)
- 4) Multiply the cosine of the angle in the

As previously discussed the magnitude of the spatial vector measured by the X and Y coordinates can be determined from the frontal plane. The magnitude projected on Z coordinate can be determined by multiplying the sine of the angle in the transverse plane by the magnitude along X coordinate (sine of angle a = cosine of angle b) (Fig 55). The magnitude of the spatial vector can be calculated. It has been demonstrated that the X Y Z coordinate values can be obtained. The magnitude of the spatial vector measured by the coordinates is determined

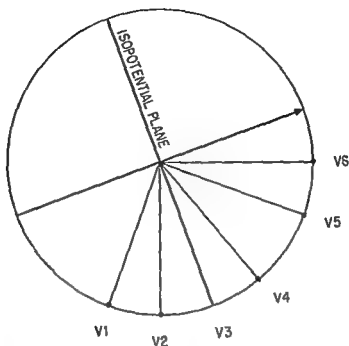


Figure 56

by utilizing the rule of squared coordinates. The magnitude of any force squared equals the sum of the squares of its projection on each of its coordinates.

Lead measurement of spatial vector magnitude = $X^2 + Y^2 + Z^2$

The calculations of magnitude given above are based on Lead I measurement of magnitude. Lead I was shown to measure 3L r of a vector. Thus the relative magnitude of the spatial vector at its point of origin in the volume conductor is lead measurement of spatial vector magni-

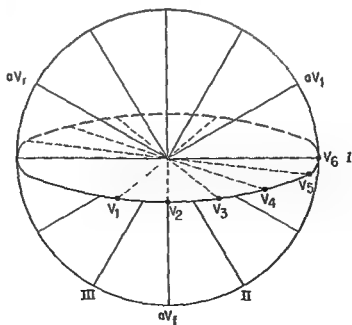


Figure 57

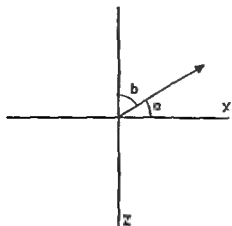


Figure 58

tude times $\frac{1}{\sin \alpha}$ where r is the radius and L is the length of Lead I

SAGITTAL PLANE

The sagittal plane is comprised of the Y and Z coordinates or V1 and V2. It is the profile of the body (Fig 59). The direction and magnitude of any vector can be determined in the

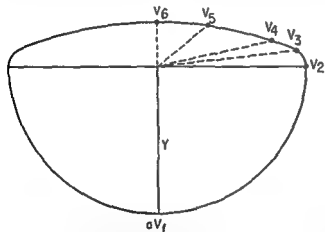


Figure 59 A right profile of the Sagittal Plane

sagittal plane by using the values obtained for Y and Z coordinates in the frontal and transverse planes

EFFECTS OF ECCENTRIC ZERO CENTER

The simplified presentation of the centric dipole concept given above is not exactly correct. The zero center is eccentric in location and the

lead lengths are not equal. The routine ECG finds its greatest application in the measurement of duration, sequence of events and relative direction of one spatial vector to another. Errors in directional measurements are largely the same for all vectors in the same individual; thus measuring the spatial angle between two vectors is practical. The normal values obtained for ECG measurements are based on the centric dipole concept and comparisons cannot be made with any other concept until other normal values are obtained. The errors imposed by the assumption of a centric dipole should be understood to evaluate the unusual circumstance and to comprehend the advances occurring in this rapidly changing field.

An example of the errors imposed can be obtained from considering the fundamentals of Lead I (Fig 60). Due to eccentric location of the zero center, Lead I is not a balanced bi-

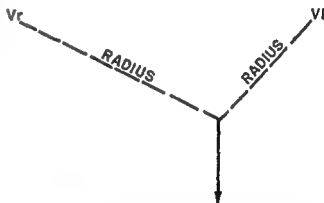


Figure 60

polar lead. The leftward position of the zero center permits the vertical and sagittal component of the spatial vector to influence the lead measurement. The vertical component directed towards the foot has a negative influence on the lead measurement. The magnitude of the influence depends upon the magnitude of the vertical component and the degree of eccentricity.

It is clear that if Lead I is to be used to measure only the X component of a spatial force, it must be a true bipolar lead. Similar effects occur in Lead II and Lead III. Usually elec-

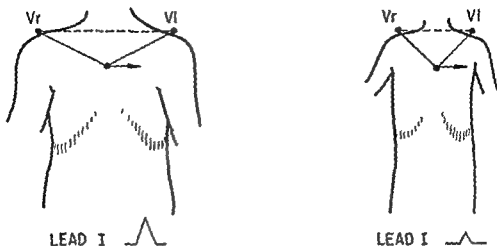


Figure 61

tude VI records a much larger negative influence from the vertical vector than Vr because the radius distance of VI is shorter and the obtuse angle is greater. Eccentricity can cause an initial negative wave in Lead I. Such factors are responsible for many ECG patterns that have been ascribed to septal depolarization and other anatomic correlations. No attempt at anatomic correlation should be made unless the leads used are true equal bipolar leads or true unipolar leads.

The V leads are also affected by eccentricity of the zero center. The value measured at the negative electrode is not really zero in these circumstances and the positive V lead electrode does not represent the point potential at its location but the difference in potential at that point and the value of the negative terminal.

LEAD LENGTH AND VECTOR AXIS

Lead length depends upon the physical contour and shape of the human body. The individual with a short squat chest has a long axis for Lead I and records a large potential value for the transverse (X) component. The individual with the long narrow chest has a relatively short Lead I axis and records a small potential value for the transverse component. The latter ECG records a more vertical vector and the former a more horizontal vector even though the actual direction of the spatial vector may be identical in both individuals. Thus the

vertical vector and transverse vector as described by the usual ECG is also an expression of the shape of the volume conductor (Fig 61).

The principle of lead length is clearly seen from the formula for magnitude. Lead I = X component magnitude times $\frac{L}{r}$. When the X component magnitude and the radius are constant the magnitude measured by Lead I is proportional to the lead length.

LIMITATIONS OF ROUTINE ELECTROCARDIOGRAPHY

The limitations in the use of the routine ECG may be summed up as follows:

- 1) Variations due to eccentric dipole position
- 2) Variations in lead length due to physical shape of the volume conductor
- 3) Variations imposed by nonhomogeneity of the volume conductor
- 4) Inability to measure the magnitude of component forces and their duration from a coordinate graph as these are measured only from the peripheral arc of the rotating central resultant
- 5) When using the direct writing instrument sensitivity to more rapid events is lost

These criticisms do not mean that routine electrocardiography is not a useful and valid tool but serve to point up that its full application has not been realized. Direct writing instruments offer the distinct advantage of a practical bedside or field instrument but they are not

adequate for study of more complicated problems or for investigation. The problems associated with the first four limitations given above are somewhat mitigated by the use of a less sensitive instrument.

The routine ECG should be used to measure duration, sequence and relative direction of spatial forces. It may be used in a somewhat more coarse fashion for magnitude measurements.

VII

The Normal Electrocardiogram

THE P WAVE

THE FIRST EVENT of the normal cardiac cycle is atrial excitation creating a P loop. The projection of this event on a lead axis creates the initial wave called the P wave. The character of the P wave in a given lead depends upon the P loop and the lead axis (Fig 62). Usually the P loop is located between zero and ninety degrees (between the axis of Lead I and aVF). The P loop is directed towards the positive pole of Lead I and II creating a positive P wave in

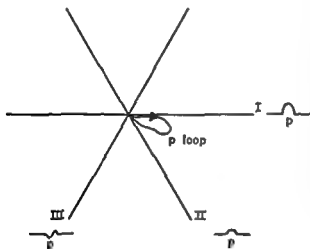


Figure 62

those leads. If the force is directed leftward parallel to Lead I, the P wave will be positive in aVI and negative in Lead III. A P loop directed downward towards aVF will cause a negative P wave in aVI. Often there is a biphasic P wave at V1 or V2 with an initial positive displacement and terminal negative component. The force of atrial excitation is above the electrode at its onset and below it as well

as posterior terminally. By more remote electrodes (V9) it is seen that the P loop is chiefly posterior. Variations in configuration are numerous.

The duration of the P wave reflects the time required for atrial excitation. Atrial excitation can require no less time than the longest P wave duration in any lead. A more precise measurement requires simultaneous recording of three mutually perpendicular leads. *Normally the P wave duration does not exceed 0.12 sec.* In general, the greater the P wave duration, the greater is the surface area of the atria.

The amplitude of the P wave is dependent upon the effective area of the wave front of excitation. In turn, this is dependent upon wall thickness. Amplitude measurements are fraught with pitfalls due to such factors as conduction and distance. For this reason, P wave amplitude is best evaluated by comparing its amplitude to other events: ventricular excitation and recovery. *A very general rule is to regard any P wave of 3 mm amplitude as unusually large; then compare to see if all the events are increased in amplitude or only the P wave.*

THE PR INTERVAL

The interval between the onset of the P wave and the onset of ventricular excitation is called the PR interval (Fig 63). It measures the time between the onset of atrial excitation and beginning of ventricular excitation. It includes the time required for atrial excitation (P wave) and the time required for transmission of the impulse through the AV node and its ramifications to ventricular muscle reception areas. The latter is represented by a nearly isoelectric interval

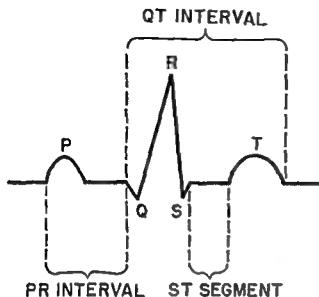


Figure 63

between the P wave and the onset of ventricular excitation (approximately 0.06 to 0.08 sec.)

The PR interval varies with age and heart rate. In the adult with a heart rate of 80 per minute the PR interval should not exceed 20 sec. A PR interval of 20 sec. at a heart rate of 90 per minute is abnormal. A PR interval of 10 sec. or less is also abnormal indicating a change in the usual order of excitation.

Care should be taken to measure the PR interval in a lead with the longest QRS duration and a sharply defined P wave of greatest duration. Otherwise the measurement may not reflect the true time required for transmission of the impulse from the SA node to ventricular muscle. Simultaneous leads obviate this difficulty.

THE QRS COMPLEX

Configuration. The forces of ventricular excitation graphed on a lead axis are called the QRS complex. By convention when the initial wave is negative it is called a Q wave. The first positive wave is called an R wave. The first negative wave (below the base line) following a positive wave is called an S wave. A positive wave following an S wave is called an R wave. Thus the initial events may create a Q wave in one lead and an R wave in another lead. A deflection of an II wave toward the base line fol-

lowed by a secondary increase in amplitude is not an S wave. Such a change is notching of the R wave.

If a wave is of small amplitude it is often printed in small letters and if large printed in capital letters. A small q wave, a large R wave and a small s wave is written qRs. When the entire QRS complex is a large negative deflection it may be called a QS deflection.

The QRS configuration in any lead depends upon the relation of the lead axis to the spatial pathway of ventricular excitation. Consider a simple pathway directed anteriorly to the left and downward then returning posteriorly to the right and to the center of origin (Fig. 64).

Lead I will have an initial R wave. The max-

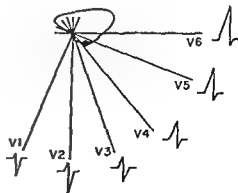
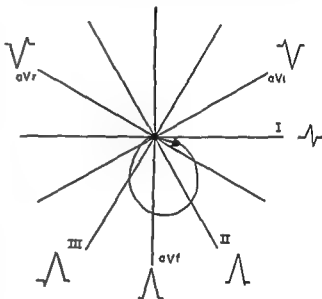


Figure 64

imum height of the R wave corresponds in time to the most leftward point on the QRS pathway. Its amplitude is proportional to the X component of the instantaneous vector occurring at that time interval. The R wave diminishes in amplitude as the pathway becomes more perpendicular to the axis of Lead I. The S wave begins as the pathway moves more than 90° away from the positive terminal of Lead I. The most negative point of the S wave corresponds to the time the spatial pathway reaches its most rightward point. The S wave diminishes as the pathway returns to the center of origin.

By remembering the lead axis the instantaneous vectors at each time interval can be graphed as a coordinate graph. Thus the QRS complex can be built for each lead.

The maximum left and right course is depicted by Lead I. The maximum downward point is the height of the R wave in aVF. By correlating simultaneous leads the spatial pathway can be reconstructed in a general manner from the QRS complexes. To accurately reflect the spatial pathway other factors such as lead length, radius distance and zero center location must be considered.

The precordial leads in a somewhat more general fashion correspond to the spatial QRS pathway. The time of the height of the R wave in V2 corresponds to the time of the maximum anterior course of the QRS pathway. The time and magnitude of the height of the R wave increases at more leftward positions. This reflects more of the spatial pathway being in the positive zone of the lead axis. The R wave in Lead V6 records the maximum leftward course of the QRS pathway. It begins to lose amplitude as the pathway loses its leftward location. The maximum posterior course of the loop corresponds to the maximum negative point (S wave) in V2.

The Intrinsicoid Deflection An electrode placed directly on a segment of heart muscle will record a positive wave as excitation moves outward to the surface. When excitation reaches the surface a deflection below the base line occurs instantly. This deflection is called the *in-*

trinsic deflection. The time between the onset of excitation and the onset of the intrinsic deflection is an index of the thickness of the muscle directly beneath the electrode. The time of onset of the intrinsic deflection is delayed when the muscle is abnormally thick.

The principle of the intrinsic deflection is indeed valid for electrodes placed directly on the heart. However, electrodes from the body surface behave differently and are in certain respects remote reflecting the action of the heart muscle as a whole. One school of investigators chose to consider the electrodes on the chest as behaving like they were influenced chiefly by the segment of muscle beneath them. Thus the downward deflection at any electrode position represented the thickness of the muscle mass beneath. This deflection being intrinsic like it was called the *intrinsicoid deflection* (oid means like).

The intrinsicoid deflection at V1 has been called an index of right ventricular wall thickness and its upper limit of normal given as 0.05 sec. The intrinsicoid deflection at V6 has been called an index of left ventricular wall thickness with a normal value up to 0.45 sec. The times are correctly measured as the onset of the deflection, i.e. the end of the peak of the R wave.

In practical application I have found these measurements of little value. When they are truly abnormal there is no difficulty in diagnosing the disorder without this determination. Moreover the chest electrodes are markedly influenced by the entire heart. The time of the R wave peak in V2 corresponds to the time of the most anterior course of spatial QRS pathway and is exactly equivalent to the time of the depth of the Q wave from an electrode directly opposite V2 on the back (Fig. 65). Thus the time interval is the same whether the electrode is on the front of the chest over the right ventricle or over the back over the left ventricle.

The peak of the S wave in V2 corresponds to the most posterior course of the spatial pathway.

This was originally defined for the intrinsic deflection measurements by Lewis T. and Reilly M. A. *Phil. Tr. R. Soc. Lond. Ser. B* 181, 1915.

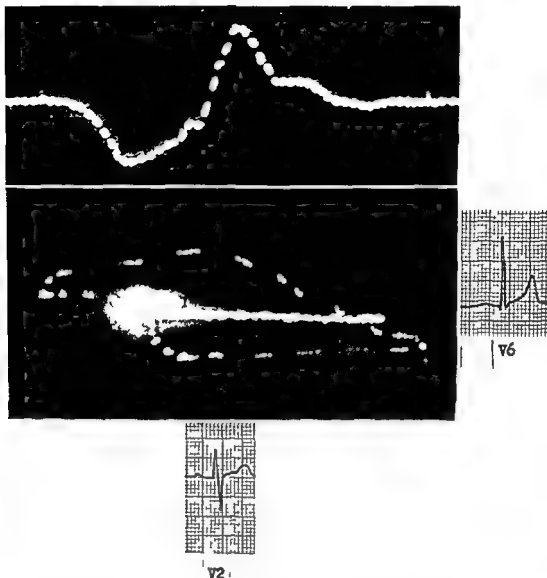


Figure 65 The interior course of the QRS loop creates the initial Q wave in the V lead recorded from the back (top of figure) and an initial R wave in V2. As the loop reaches its most leftward point the height of the R wave in V6 is recorded. The terminal rightward course of the loop creates the terminal S wave at V6. Minor variations in initial Q waves and R waves are often due to placing the chest electrodes above or below the zero center.

When this occurs they are no longer strictly recording the transverse plane.

The height of the R wave (intrinsicoid deflection) recorded by an electrode over the back occurs at the same time as the depth of the S wave at V2. Thus the "intrinsicoid time of the left ventricle" can be obtained from an electrode very proximal to the right ventricle. These observations are not compatible with the notion that chest electrodes behave like proximal electrodes in reflecting chiefly the activity of the muscle segment beneath them.

The intrinsicoid deflection at V6 actually measures the time of the most leftward course of the spatial pathway. Since the left ventricle is posterior and the septum is anterior, much of left ventricular activation creates a posterior force occurring after the intrinsicoid deflection at V6. In the routine ECG left ventricular activation is often best represented by the S wave of V2.

Since the magnitude of the leftward course

normally increases as the right ventricular force declines the maximum leftward magnitude occurs more closely to completion of right ventricular activity. For this reason the height of the R wave at V6 closely corresponds to the completion of right ventricular activation in the normal heart rather than the left ventricle. Indeed the time commonly given for completion of right ventricle activation (0.03 to 0.04 sec) will be found in the normal heart to be the time for the height of the R wave at V6.*

Duration The QRS duration is measured to determine the length of time required for ventricular excitation. The longest QRS duration in any lead approximates the excitation interval. Lead V2 most often has the greatest QRS duration. The terminal activation of the left ventricle creates a posterior pathway. The ideal determination is made from three simultaneous mutually perpendicular leads. The QRS duration is an index of wall thickness. Since the wall becomes thicker from childhood to maturity the QRS duration increases accordingly. The normal QRS duration varies from 0.06 to 0.10 sec.

Due to complications of the wave front at the base of the heart discussed previously the terminal QRS complex may be complicated or prolonged. This is particularly true in younger individuals. With maturity these variations are less common.

Amplitude QRS amplitude is dependent upon the recording technique (lead lengths, electrode distances), density of charge across the wave front and the volume of the ventricles. The volume of the ventricles depends upon cardiac rate and phase of the respiratory cycle. Amplitude should be determined by using three perpendicular leads. A rough guide for QRS amplitude is the sum of the QRS amplitude in V2, aVF and V6. When this value exceeds 50 mv, increased amplitude should be considered.

The right ventricle begins activation 0.05 sec after the left ventricle. By 0.02 sec the confluent right ventricular cone is established. Muscle spread out and outward at 3 mm each 0.01 sec. Allowing a wall thickness of 6 mm, the right ventricle will complete activation by 0.04 sec.

Amplitude measurements of spatial forces require consideration of three perpendicular axes.

Effects of Heart Rate With a slow heart rate the stroke volume is increased. The increased volume is reflected in increased QRS amplitude. A rapid heart rate is associated with smaller QRS amplitude (Fig. 66).

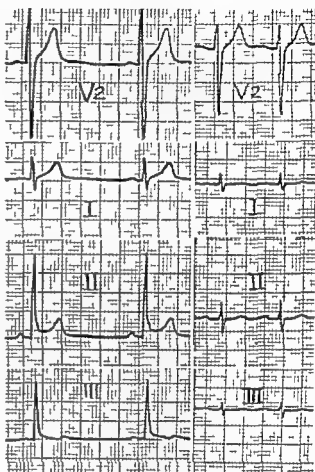


Figure 66 Note the difference in amplitude in two individuals of approximately the same age, health and size. Increased amplitude is seen with a rate of 55 per minute and low amplitude at a rate of 100 per minute.

Effects of Respiration Respiration has a large effect on QRS amplitude and configuration. In the past these have been attributed to a simple change in heart position. An original observation indicates that the change is not a simple axis shift. In actuality the ECG is a sensitive measurement in the changes in left ventricular stroke volume. It is well known that inspiration

causes increased right ventricular stroke volume and diminishing left ventricular stroke volume.

During deep inspiration there is a progressive decrease in QRS amplitude in all leads. The initial 0.4 sec period becomes more vertical with increased right ventricular volume. The terminal 0.4 sec is diminished consistent with decreased left ventricular volume. A simple shift in axis requires that amplitude be increased in Lead II (Figs 67 and 68). Holding the

amplitude and increase in size (Fig 71). This may be observed at quiet respiration in individuals with adequate stroke volume. It will be recalled that the S wave of V2 is chiefly due to the left ventricle.

The influence of respiration is further complicated by the increase in cardiac rate during inspiration. The increased rate can decrease diastolic filling with decreased volume causing diminished amplitude. However when the rate



Figure 67 A simultaneous recording of four leads showing the decrease in QRS amplitude with inspiration and return to normal amplitude with expiration. Note the principal effect is on the terminal half of the QRS complex not the first half. The last half of the QRS cycle is largely created by the left ventricle.

breath at deep inspiration causes slowing of the heart rate. In the vertical electrical axis it is noted that an S wave in Lead I disappears or is markedly diminished. A simple right axis shift would have the opposite effect increasing the S wave (Fig 69).

With inspiration a late R wave in aVR will lose its prominence and a deep S wave may occur in aVL. This terminal change in the QRS events simply reflects a leftward (more vertical) shift of the terminal QRS forces. A shift in rightward axis with inspiration of necessity would increase the late R wave in aVR (Fig 70).

A most striking example of the influence of respiration is noted in V2. The S wave behaves as a simple spirogram by its progressive dim-

inution and increase in size (Fig 71). This may be observed at quiet respiration in individuals with adequate stroke volume. It will be recalled that the S wave of V2 is chiefly due to the left ventricle.

Change in conduction with lung inflation can not be used as an explanation of diminished amplitude as the initial 0.4 sec show increased amplitude. Moreover V2 position is directly over the heart and not insulated by lung tissue even in full inspiration. The drop in QRS amplitude is also manifested in all leads when a premature nodal or atrial contraction occurs early enough to markedly diminish diastolic filling.

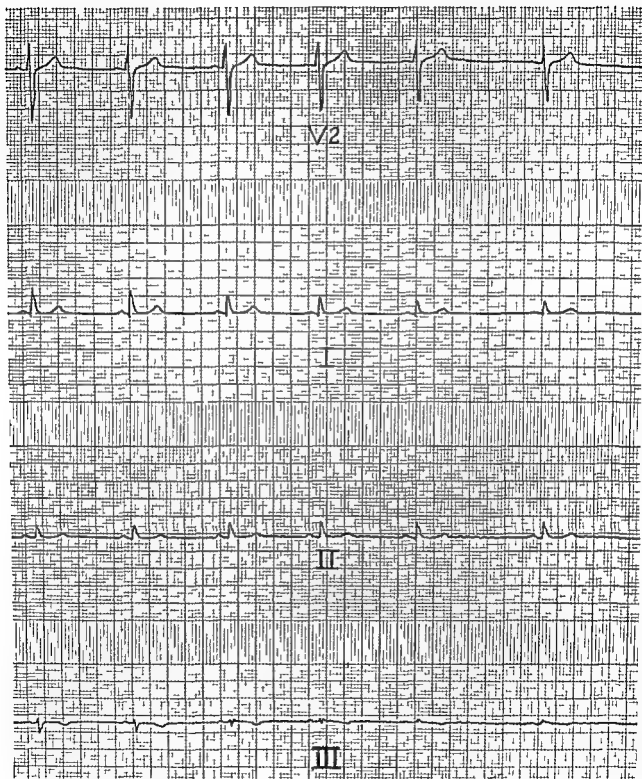


Figure 68 Another example of the overall change in QRS amplitude caused by inspiration. A simple anatomic change in cardiac position due to the lowering of the diaphragm should cause increased amplitude in Lead II as the electrical position becomes more vertical. This simply does not occur.

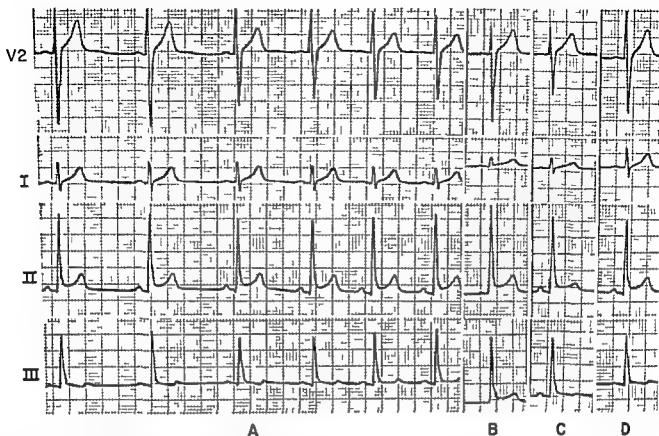


Figure 69 A) The inspiratory phase of the respiratory cycle with decreased amplitude of the S wave in V2 B) Holding the breath at the height of inspiration. Note the S wave in Lead I entirely disappears completely contrary to a simple right axis shift C) After holding the breath for 30 seconds at full inspiration the QRS amplitude increases despite the fact that insulation and mechanical effects of lung inflation are at the maximum D) On expiration about the fifth cycle later the QRS amplitude is the same as at the onset of the maneuver

This is not associated with respiration (Fig 72). Also against the theory of insulation is increased P loop magnitude observed during inspiration and decreased P loop magnitude with expiration. Moreover insulation is theoretically greatest with deep inspiration and with prolonged held deep inspiration the ECG changes back toward normal.

The observation that the ECG can be used as an index of difference in right and left ventricular stroke volume creates a whole new avenue of investigation in electrocardiography including the possibilities of application to pulmonary function. It is interesting to note that the ballistocardiograph has long been touted as superior to the ECG because it could be correlated to stroke volume. Changes in the ballistocardiograph have been correlated to the

respiratory cycle. The above observations indicate the ECG may be useful for this measurement.

Effects of Exercise The usual response of the young lean individual to acute exercise is an increase in rate maintaining stroke volume. The QRS amplitude remains unchanged. There are a group of individuals that respond differently. Immediately after exercise they have a decrease in QRS amplitude associated with an increase in rate. As the rate slows the QRS amplitude returns to normal. Although the group observed to date is small it appears to be in those individuals who are overweight and do little physical exercise. One explanation for their decreased QRS amplitude is diminished diastolic filling associated with rate and inadequate venous return (Figs 73, 74 and 75).

THE NORMAL ST SEGMENT

After the QRS complex the phase of ventricular recovery begins. The early stage of recovery creates very small forces due to the size

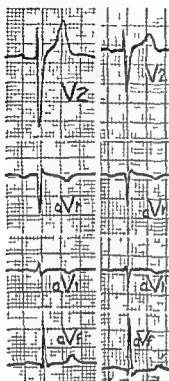


Figure 70 Note the small decrease in the terminal R wave in aVR from expiration (left) to inspiration (right)

and shape of the wave front. This early period is represented by a nearly isoelectric phase on the electrocardiogram and is called the ST segment. The initial anterior direction of the small forces originating from the right ventricle creates a normal elevation of the ST segment at V2. The proximity of the electrode at V2 position enhances this effect. Particularly in young individuals normal ST segment elevations may be quite marked (Fig 76). A slow cardiac rate is commonly associated with this finding. Normal ST segment elevation occurs in the leads with a positive wave for the entire event of ventricular recovery (leads with an upright T wave).

THE NORMAL T WAVE

Recovery of the left ventricle from apex to base creates a much larger force manifested by the T wave. In general the T wave is positive in those leads with positive QRS complexes. Normal variations exist due to the difference in size, shape, and speed in the wave front of recovery and excitation. The duration of the T wave reflects the time required for recovery. The main T wave deflection occupies roughly 16 sec. The entire recovery time from completion of QRS to completion of T approximates 28 to 36 sec. The T wave amplitude is greatest in the lead most nearly parallel to the greatest displacement of the T loop (allowing for electrode distance).

In the adult the T wave is normally upright in leads I, II, and precordial leads V3 through V6. Depending upon electrical axis and lead length the T wave may be either positive, negative, or isoelectric in all other leads except aVR. Lead aVR normally has a negative T wave in the adult.

Amplitude of the T wave is very difficult to define in precise figures. In young individuals the T wave may be extraordinarily large. This is particularly true of the more athletic individual with a slow heart rate. In general a T wave of seven tenths millivolts (7 mm at normal standard) is large, requiring consideration of age, rate, electrode distance, and physiological stages.

The T wave is normally subject to variability. Drinking ice water to chill the apex reverses the apex base order of recovery, inverting the T waves in those leads with positive electrodes facing the apex. Changes in cardiac filling due to change in cardiac rate or changes in posture may also change the characteristic of the T wave.

The African ECG is particularly variable in reference to transitory T wave changes. A young adult negro with no demonstrable heart disease often presents a seeming abnormal record with a high degree of variability. This is thought to

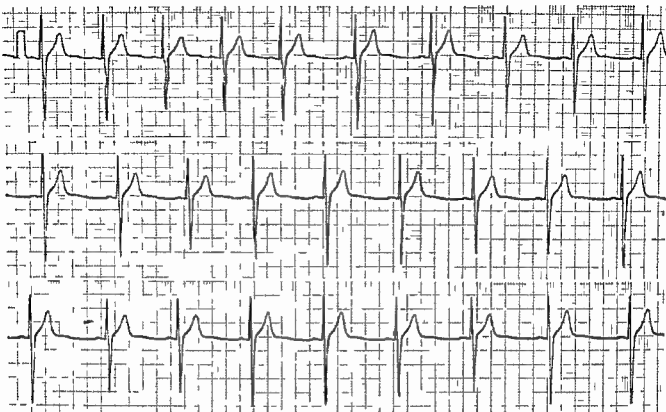


Figure 71 This is a continuous strip of Lead V2 showing the perfect Sprogram recorded by the S wave. Note there is no change in the R wave (first half of the QRS) and the base line is absolutely stable. This is a selective change of the S wave alone.

occur more frequently during the time the juvenile pattern is changing to the adult characteristics (Fig 77).

THE NORMAL U WAVE

The after potential force is demonstrated by a small positive wave immediately following the positive T wave. Little factual information regarding its normal and abnormal variations is available (Fig 78).

THE NORMAL QT INTERVAL

The interval between the onset of ventricular excitation and the end of ventricular recovery is called the QT interval. This period of time corresponds approximately to mechanical systole of the heart and is sometimes called electrical systole. The QT interval is properly measured as the longest interval in any of the leads. Its value will vary with the heart rate.

The QT interval should be corrected for rate before determining whether it is within normal limits. This may be done by utilizing Bazette's formula:

$$K = \text{QT interval} \times \sqrt{R - R \text{ interval}}$$

K is a constant and is .37 sec for men and .40 sec for women. $R - R$ interval means the time between two successive R waves. The QT interval is measured directly from the tracing in question. Whenever QT interval times $\sqrt{R - R}$ is greater than the constant K , the QT interval is prolonged. An average QT interval is .34 to .44 sec.

THE MEAN VECTOR

A number of normal values in electrocardiography are dependent upon the so-called "mean vector." These are derived from the net positive or negative value of complexes graphed on the lead coordinates. The QRS complex as an example will have a net positive, negative, or zero

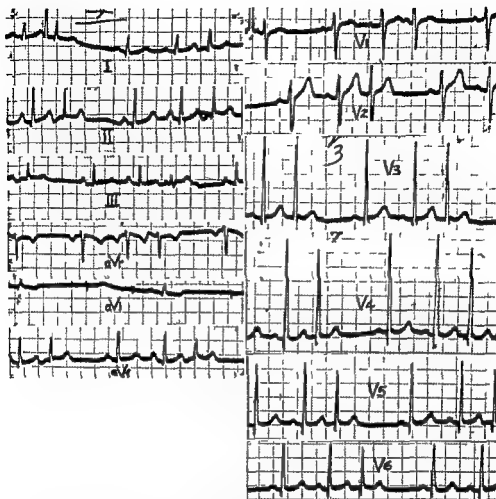


Figure 72 Each nodal premature contraction has less QRS amplitude than the preceding and following QRS complex. This appears to occur only with very early premature contractions.

value in any lead. The same is true for the other events. The final mean vector determined for an event is an overall resultant force equated to the event.

The first step is to determine the net value of a complex in a given lead. The triangular area of the positive and negative deflection is obtained (amplitude times base divided by two). The values are subtracted (added algebraically) to obtain the net value. When the positive and negative deflections are equal the net value is zero. A lead with such a complex is called the isoelectrical lead for the event being measured. The mean vector is perpendicular to this lead. Given Lead I with a diphasic QRS complex and net value of zero, the mean QRS axis is perpen-

dicular to Lead I. If Lead II and III are positive the mean QRS axis is $+90^\circ$. Determination of the mean T and P vectors may be made in a similar manner.

The mean vector measurements are clearly mathematical derivatives and crude determinations. The reader should review again the mathematical origin of mean vectors obtained from coordinate graphs as discussed in Chapter I. They can be used for average directions and in a very coarse manner for magnitude.

THE MEAN QRS VECTOR

On occasion it is advantageous to calculate the mean vector acting for a specific period of time. The mean QRS vector acting during the

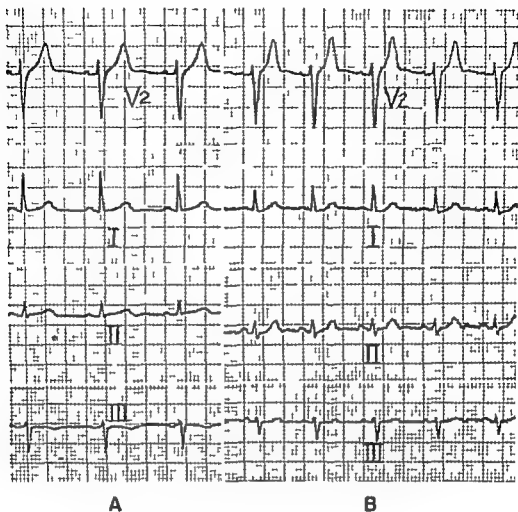


Figure 73 A) Before exercise B) Immediately after exercise

initial 0.4 sec of ventricular excitation is manifested only during this 0.4 sec interval irrespective of the character of the QRS complex during the latter period of excitation. The terminal 0.4 sec QRS vector acts only during the last 0.4 sec of ventricular excitation.

The relationship of these component QRS vectors to the mean QRS vector may be illustrated by the following case (Fig 79).

1) The total QRS duration is 0.8 sec (longest QRS duration in any lead).

2) Lead III has a QRS duration of 0.8 sec. Its initial 0.4 sec is a Q wave 4 mm deep. The last 0.4 sec is an R wave 4 mm high. The initial 0.4 QRS value is minus 2 units (4×1 divided by 2) and its terminal QRS value is plus 2 (4×1 divided by 2). The mean QRS value in Lead III then is 0.

3) Lead I has a QRS duration of only 0.4 sec. The complex has 8 mm of positive amplitude. The mean QRS value in Lead I is 4 positive units (8×1 divided by 2). Since the QRS interval was short in Lead I, an isoelectric interval of 0.4 sec duration must have occurred on this lead axis during ventricular excitation. This occurred either before or after the positive complex was inscribed.

4) Lead II has a QRS duration of 0.8 sec. The entire complex is positive. Its height of 4 mm is reached at 0.4 sec. The initial 0.4 sec QRS value is plus 2 units (4×1 divided by 2). The terminal 0.4 QRS value is also plus 2 units. The mean QRS value in Lead II is plus 4 units (4×2 divided by 2).

The initial 0.4 QRS value changes from minus 2 units in Lead III to plus 2 units in Lead I.

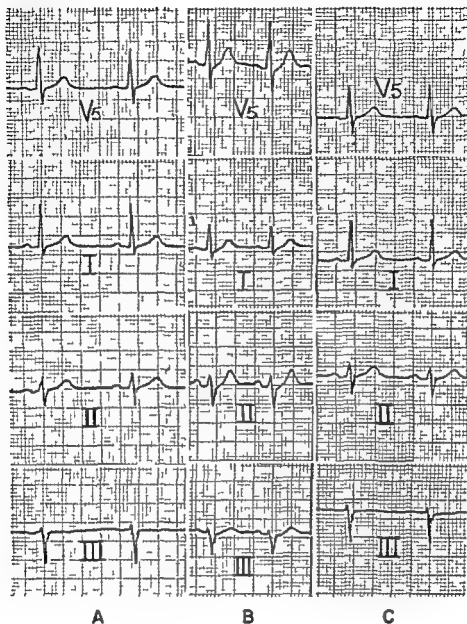


Figure 74 A) Before exercise B) Immediately after exercise C) Return to normal rate after exercise

Therefore an isoelectric point must exist between Lead I and Lead III for the initial 0.04 sec vector. This isoelectric point was at Lead II and wrote an isoelectric interval before the positive wave occurred. This also caused the QRS interval to appear shortened in Lead II (0.04 sec). The initial 0.04 sec mean QRS vector is perpendicular to the lead axis of Lead II and directed toward Lead I or its axis is -30° . The terminal 0.04 sec mean QRS vector records plus 2

units in Lead III plus 2 units in Lead I and plus 4 units in Lead II. Its axis is parallel to the axis of Lead II or $+60^\circ$. The mean QRS complex is perpendicular to Lead III where the mean QRS magnitude was 0. The mean QRS vector has an axis of $+30^\circ$.

The anterior or posterior direction of the 0.04 sec vector can be determined from the V leadsogram by locating the transition from positive to negative value.

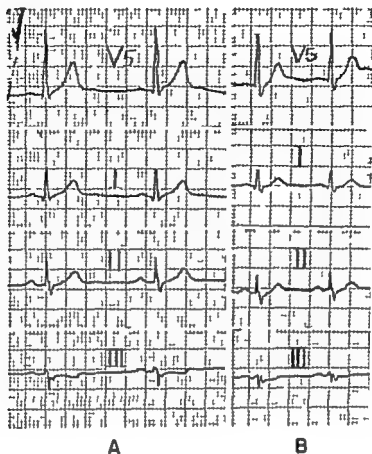


Figure 75 A) Before exercise B) Immediately after exercise

NORMAL MLAN QRS AXIS

The projection of the QRS vector on the frontal plane is called the QRS axis. It is defined in degrees from the axis of Lead I. The normal adult heart may have a QRS axis between -30° and $+110^\circ$. Any axis beyond this range must be explained before the axis is considered normal. In the infant the axis is normally to the right beyond $+90^\circ$ and gradually shifts leftward to assume the adult position. As age advances the QRS axis commonly shifts leftward to an axis of 0° or a minus value.

As the QRS vector is rotated out of the frontal plane to either a more posterior or anterior direction its magnitude in the frontal plane is diminished. In the normal adult the mean QRS vector is directed posteriorly. This is the influence of early completion of septal activation causing the terminal QRS pathway to be di-

rected posteriorly. If the left ventricle is elongated the septal content is larger and the mean QRS vector is directed even more leftward and backward.

The axis of the QRS vector in the frontal plane or any other plane can be determined by the methods outlined previously for the calculation of any spatial force in any plane.

THE MEAN T VECTOR

The spatial T vector is directed far to the right and posteriorly at birth. The vector gradually moves leftward and anteriorly with age. Occasionally the posterior direction of the T vector may persist into adult life. This is one cause of negative T waves in V1 through V3 and it is called a persistent juvenile pattern.

The normal mean T vector in the adult approximates the longitudinal axis of the left ven-

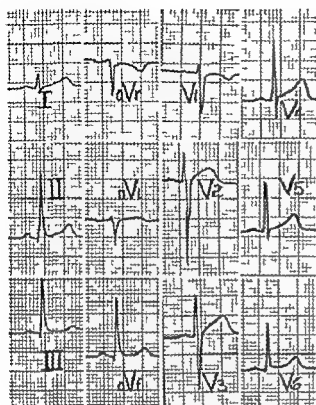


Figure 76 Note ST segment elevation in Leads II III aVF V2 and V3 in a healthy subject

tricle. Thus it is normally anterior to the mean QRS vector. When the mean QRS vector becomes markedly leftward the mean T vector will be to its right e.g. QRS -30° T vector zero degrees.

The magnitude of the normal T vector is usually less than the magnitude of the QRS vector although it may be larger in slow hearts with vertical axis as seen in young individuals.

THE MEAN SPATIAL QRS-T ANGLE

Chiefly due to early completion of septal activation a normal angle exists between the mean QRS and mean T vectors (Fig 80). The spatial angle in the adult is usually less than 60° . Occasional individuals without known evidence of heart disease may have an angle as large as 90° . These are exceptional. Any angle over 60° must be explained.

In the normal young adult the ventricles

are short. The septal component is small. The mean QRS vector tends to be electrically vertical and not very far posterior. It is chiefly straight down. The QRS and T vectors are nearly parallel and the mean spatial QRS-T angle may be only 15° . Later in life the left ventricle tends to elongate increasing the ef-

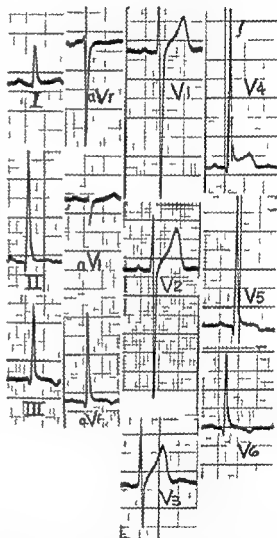


Figure 77 This ECG was recorded on a young negro without any demonstrable evidence of heart disease, hypertrophy or hypertension. The QRS amplitude is increased in part due to the small chest. Note the QRS complex being recorded off the paper. This is an error in technique. When the QRS amplitude is too large for proper recording the standardization should be reduced.

fective magnitude of the septal r vector. This causes a shift in the QRS axis leftward and posterior. The mean spatial QRS-T angle widens.

If the magnitude, direction, and sense of the QRS and T vector are known, the net positive or negative value that is recorded in any lead can be determined (Fig. 80).

If the spatial QRS-T angle is abnormally

wide, it will register abnormal patterns measured by individual leads. The measurement of the QRS-T angle projection on only one plane is fallacious and misleading. In any instance when the QRS and T vector have the same frontal plane axis, the mean QRS-T angle in the frontal plane is 0°. If the QRS vector is posterior and the T vector is anterior, the spatial angle will become smaller as it becomes vertical.

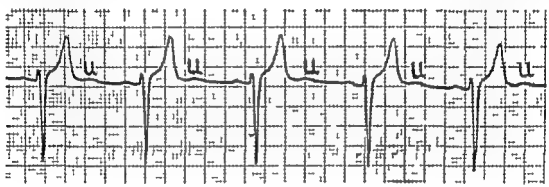


Figure 76

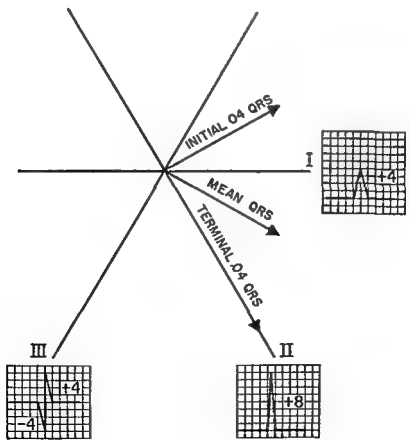
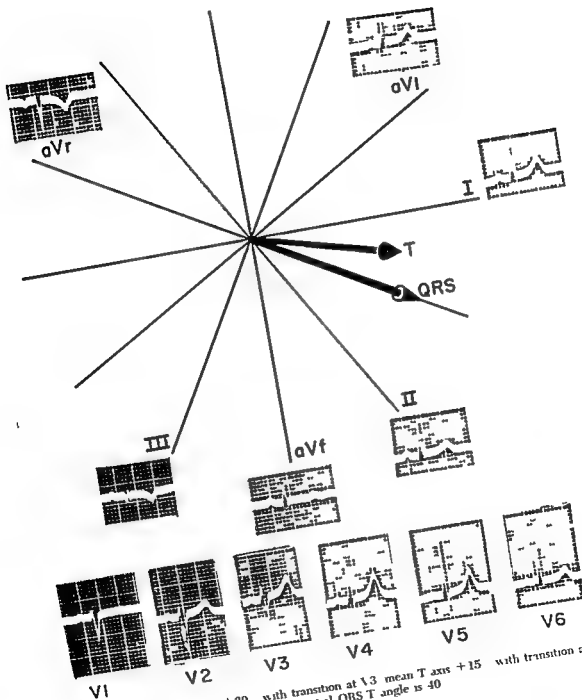


Figure 79

The Normal Electrocardiogram



(Fig 81) Viewed in the sagittal plane the angle between *a* and *b* is larger than the more vertical angle between *a* and *b* although in both cases the angle is 0 as projected on the frontal plane and 60° as projected on the transverse plane.

If the QRS and T vectors both have an axis of $+90^\circ$ the angle will be either 0 or 180° in the transverse plane projection. Further increase in the spatial angle will only increase the magnitude of projection in the transverse plane (Fig 81). This accounts for the wide QRS-T angle in the V leads in the presence of a vertical electrical axis and small mean spatial QRS-T angle.

Certain difficulties in calculating the spatial angle are inherent to the human body and must be assumed. Not only is there an individual

anatomic variation in the V lead position but the location of the cardiac zero center is not constant. The center of zero potential may be either anterior or posterior to its expected location or it may be lower or higher. When the zero center is shifted the entire reference system is altered. Nevertheless it is better to have some means of approximating the spatial angle than none at all.

The actual calculation of the spatial QRS-T angle is somewhat cumbersome and for this reason a simplified table is provided to approximate readily the spatial QRS-T angle (see spatial angle chart). Each vector is described by its axis in the frontal plane and its transition point in the V leads. The description of the vector in the left hand margin is located for either the QRS or T vector. The description of the other vector is located at the bottom of the table. At the intersection of the columns the value of the spatial angle is recorded. The method of using the table is exactly like using an ordinary index chart. The values are accurate mathematically within 5°. The chart may be used to calculate any two spatial vector forces including the angle between 04 QRS vectors and the mean QRS vector.

Assuming the lead axis of V6 is parallel to the X coordinate when a complex is transitional at V6 the vector for that event must be $+90^\circ$ or -90° in the frontal plane. Accordingly a transition complex at V6 is described in the chart as a strictly vertical force of $+90^\circ$ or -90° . If the vector is only slightly posterior V1 through V5 will be negative. If it is slightly anterior V6 is negative. The more anterior or posterior the direction of the spatial force the smaller its magnitude in lead V6.

Transition at V6 with a diphasic or isoelectric value in all the limb leads including aVF is produced by a vector which is directed entirely anteriorly or posteriorly. Such a vector is truly perpendicular to the frontal plane. If it is anterior V1 through V5 are positive. If the vector is directed posteriorly V1 through V5 are negative. On the chart they are described as anterior to the frontal plane with transition at V6.

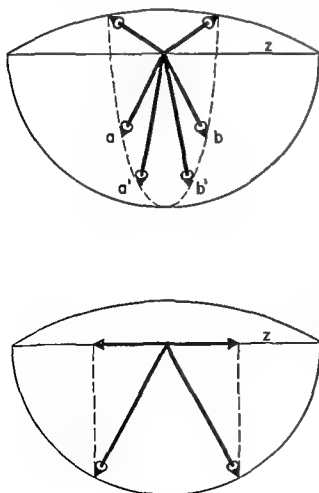


Figure 81

(A V6) or posterior to the frontal plane with transition at V6 (P V6)

In any individual case the spatial angle between two forces can be determined by using the law of cosines (Fig 82). If a line is drawn between the terminations of the QRS and T vectors a spatial triangle is formed. By the law

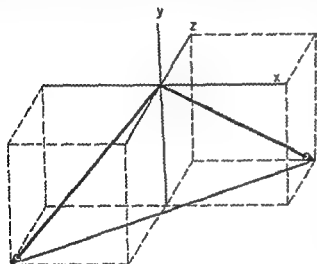


Figure 82

of cosines the angles of any triangle can be determined whenever the magnitudes of the three sides are known. One side of the triangle is the QRS vector and another side is the T vector. The base (B) must be calculated. The base can be calculated because the twelve lead ECG measures the spatial force along three perpendicular axes X, Y, and Z. The projection of the force on each axis is known for both the QRS and T vector. The sum of the square of difference of projected magnitude in each coordinate equals the square of the base. This may be expressed

$$(X_q - X_t)^2 + (Y_q - Y_t)^2 + (Z_q - Z_t)^2 = B^2$$

Utilizing the law of cosines the

$$\cosine\ of\ the\ QRS\ T\ Angle = \frac{(QRS\ vector)^2 + (T\ vector)^2 - B^2}{2 \times QRS\ vector \times T\ vector}$$

Stepwise one determines

- 1) The spatial magnitude of the QRS vector
- 2) the spatial magnitude of the T vector
- 3) the value of Base squared
- 4) and the cosine of the QRS T Angle by the law of cosines

VIII

Fundamentals of Vectorcardiography

THE PREVIOUS DISCUSSIONS have emphasized that the electrical events of the heart describe spatial pathways or loops P QRS and T. The vectorcardiogram strives to record these spatial loops. The spatial loops are polygons and the ECG records the successive resultants of the polygon. The differences and similarities of the VCG and ECG are those which exist between polygons and coordinate graphs of polygon resultants. It is clear that the ECG measures duration sequence and direction. The VCG also measures magnitude because it records the sides of the polygon is the spatial pathway.

The Frontal Plane. Vectorcardiograms are recorded with a cathode ray oscilloscope. Use is made of the X and Y axis of the orthode ray beam (side to side and up and down) created by the horizontal and vertical plates. The lead chosen to represent the X coordinate is fed into the X axis. The lead representing the Y coordinate is fed into the Y axis. When the X and Y coordinates are recorded simultaneously they record the frontal plane projection of spatial loops (Fig. 83). The six limb leads of the ECG can be extracted from this projection.

The Sagittal Plane. projection of the spatial loops requires use of the Z and Y coordinate leads. The lead representing the Z coordinate is fed into the X axis of the oscilloscope. The Y coordinate lead is fed into the Y axis. Their simultaneous recording is the sagittal plane of the spatial loops (Fig. 84).

When an oscilloscope with two beams is used the sagittal and frontal plane projections of the vectorcardiogram can be recorded simultaneously. This gives a frontal and profile view of any

one cardiac cycle. The two planes utilize all three mutually perpendicular coordinates (X, Y, Z) thus defining spatial forces.

The Transverse Plane. is recorded by feeding the lead for X coordinate into the X axis and



Figure 83

the lead for the Z coordinate into the Y axis (Fig. 85). The transverse plane represents the cross section through the thorax. The V leads may be extracted from the transverse plane by



Figure 84

from their recorded vectorcardiogram. This in no way validates these particular leads as true representatives of X Y Z coordinates for the body. In any system employing bipolar leads the leads must be centered in reference to the zero center and the leads standardized equally to each other. Standardization requires that both the lead length and the radius distance of the electrodes from the zero center be known.

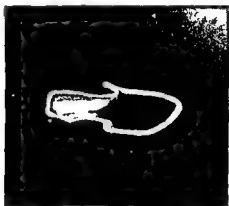


Figure 85

projecting the transverse plane upon the coordinate axis of each V lead.

Timing is accomplished by introducing interruptions in the beam at known intervals (Fig 86). If the beam is blinked out each 0.025 sec an interruption will occur in the loop every 0.025 sec. By locating the time intervals on both the frontal and sagittal planes the point in space for each instantaneous vector can be described in terms of its X Y Z coordinates. From such a determination a spatial representation of the spatial pathway (polygon) can be constructed. The instantaneous vectors of such a three dimensional representation are the successive resultants of the sides of the polygon. The successive instantaneous spatial vectors can be projected upon each lead axis to construct each lead (Fig 87).

It is obvious that whatever leads one uses for X Y Z coordinates can in turn be extracted

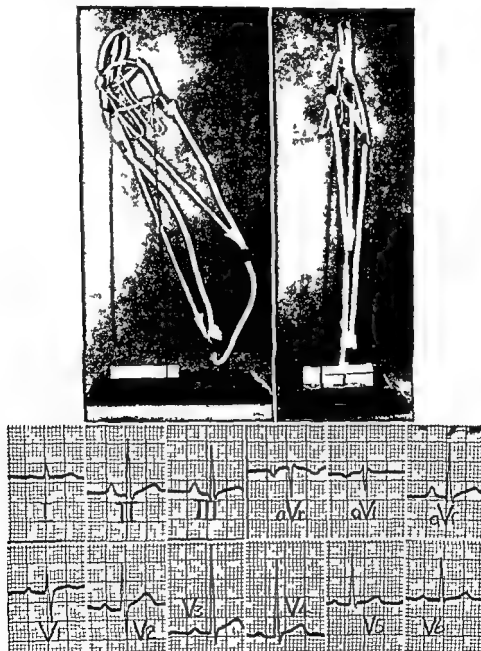


Figure 86

THE VCG SYSTEM

The system used to record the vectorcardiograms demonstrated in this text attempts to satisfy the requirements of 1) symmetrical relation to the zero center 2) measured lead lengths and 3) known radius distances of electrodes from the zero center. The first step is location of the zero center. It is assumed that the center of the left ventricle approximates this

point. A left lateral and PA chest X-ray is obtained. In each subject the center is located as a) one half the vertical distance between the right atrial groove and the left leaf of the diaphragm b) one half the distance between the mid sternal line and the left cardiac border at this level and c) one half the distance between the anterior and posterior cardiac borders at the same vertical level as located on the sagittal X-ray (Fig. 85)



The transverse lead (X coordinate) is on the back across the shoulders. It is arranged so that its length is bisected by a perpendicular extending to the cardiac center previously measured. In the adult the length most often chosen is 22 cm. Thus if the center is 4 cm to the left of the mid spinal line the left electrode (+) is 15 cm to the left of the midline and the right electrode (-) 7 cm to the right of the midline. The narrowness of the lower part of the thorax prevents symmetrical location of a lead of such length at this location in most normal adults. For this reason the lead is placed across the upper thorax. It is desirable to return lead length whenever possible.*

The vertical lead (Y coordinate) is bisected by a perpendicular from it to the cardiac center. The usual lead length in the adult is 30 cm. The negative electrode is the same as the right electrode for the transverse lead. The positive electrode is directly 30 cm beneath it.

The three symmetrically placed electrodes are all of equal radius distance from the center. The chief difference in their sensitivity is their difference in lead length. Since the vertical lead is the longest it will have the greatest sensitivity.

It is recognized that the left shoulder has a more complicated potential field. However the advantages gained by a long lead length symmetrical to the zero center outweigh this disadvantage.

In order that they be equal the standardization (amplification) must be adjusted. This is done simply by multiplying the transverse lead standardization by the ratio of the lead lengths. With a standardization of 10 cm = 1 mv for the transverse lead (length 22 cm) the vertical lead (30 cm) standardization should be 73 cm = 1 mv.

$$100 \times \frac{22}{30} = 73$$

The sagittal lead (Z coordinate) is a V lead placed directly over the cardiac center on the back (commonly 4 cm to the left of the mid spine). The standard Wilson central terminal is utilized. The lead length is obtained by measuring the distance on the sagittal X ray from the center to the back (commonly 19 cm). The standardization is determined by using the equation for the sensitivity of the bipolar lead ($3L/R^*$) and the V lead ($3/r$). When the standardization for the transverse lead is 10 cm = 1 mv (transverse lead 22 cm long, vertical lead 30 cm long and sagittal lead 19 cm long) the formula is

$$V \text{ lead standardization} = \frac{r \times \text{transverse lead length}}{R}$$

$$42 = \frac{19 \times 19 \times 22}{18500}$$

Like all systems leading from the human body, certain errors must be assumed. The indi-

Figure 87. The three dimensional VCG model is constructed from a simultaneously recorded frontal and sagittal plane VCG (frontal view at the left and sagittal view at the right). The very large white loop is the QRS loop. The black markers represent 0.1 sec intervals. The white vectors are the instantaneous spatial vectors at each 0.1 sec (the resultants of the polygon). The QRS loop begins anteriorly and sweeps downward to the left and posterior. Its terminal portion is complicated reflecting the terminal complications of the wave front. The sagittal presentation is a left profile of the model. The small P loop in this instance is more vertical than the more leftward T loop.

The white marker at the base of the model represents 5 cm length and is a standardization. By comparing the size of the standard in each succeeding photograph a concept of relative size can be appreciated. All succeeding models, unless otherwise stated, are built to the same scale in terms of electrical moment (corrected for lead length and distance of electrode positions from the heart). Each centimeter equals 5 millivolts as electrical moment.

The above example depicts a vertical QRS loop. Note the initial Q wave in Lead II, III, and aVF as the terminal QRS pathway is directed even farther toward the right shoulder. This event also creates a terminal R wave in aVR and aVL. As the terminal QRS pathway is nearly perpendicular to the axis of Lead I it has almost no influence on that lead.

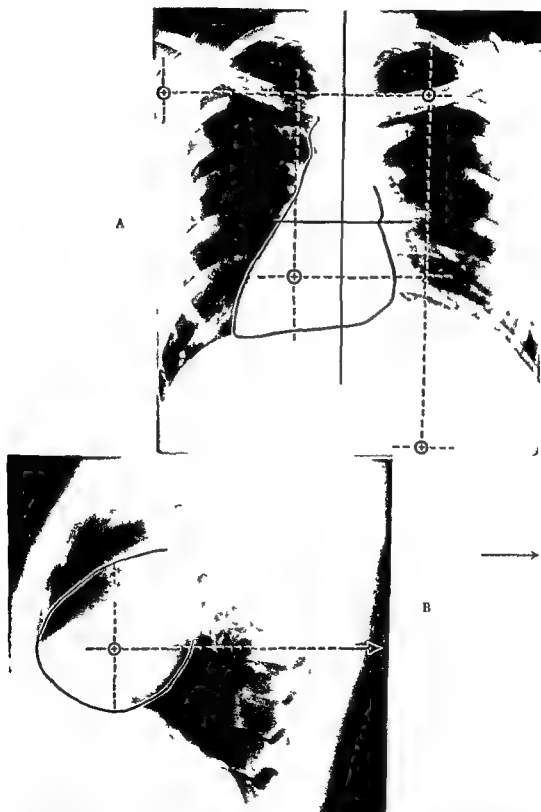


Figure 88 A) Demonstration of electrode positions in relation to the center of the left ventricle as seen on the PA film B) Demonstration of the method of determining the sagittal lead length from the sagittal x ray C) Diagram of actual electrode placement for recording VCGs

vidual variation in body shape, body size, and the location of the zero center are in part corrected in this system. It also enables conversion for factors of radius distance and lead length from one subject to the next (terms of electrical moment) which cannot be done unless such measurements are made. Without such conversion factors electrocardiographic and vectorcardiographic measurements of magnitude are invalid. Unless the leads are standardized in relation to each other, the loops are distorted.

VECTORCARDIOGRAPH ELECTRODE PLACEMENT

ELECTRICAL CENTER
OF HEART

⊗ ELECTRODE
● WILSON JUNCTION



Figure 88c

The general configuration of the complexes of P, QRS, T in the 12 lead routine ECG leads can be extracted from the VCC. Simply draw the lead axis through the center of the vectorcardiogram and graph the instantaneous vectors on the lead axis like a simple coordinate graph. Differences between the extracted leads and recorded leads can be attributed to difference in the lead systems employed. The routine ECG makes no corrections for zero center location, lead lengths or radius distance.

INDEX OF MAXIMUM POTENTIAL AND POTENTIAL SECONDS

The vectorcardiogram enables the calculation of the *Index of Maximum Potential* and the *Index of Potential Seconds*. The former is directly related to left ventricular surface area (and consequently stroke volume) and the latter is an expression of wall thickness as well. The index of maximum potential is obtained for the left ventricle simply by measuring the length of the QRS pathway after completion of right ventricular and septal activation. The pathway represents the length of the sides of the polygon.

The instantaneous long left ventricular vector commonly represents the resultant of the base vector and septal vector. Thus the last half of the QRS has two major directions: 1) a longitudinal and 2) a posterior component. The longitudinal component represents the area of the base and the posterior component represents the area of the septum. These two components are roughly two sides of a triangle (semiparallelogram) and the long left ventricle vector is the diagonal or their resultant.

The index of potential seconds is obtained by first multiplying the length of each segment of the spatial pathway by its duration of action, then summing these values. Each segment represents a side of the polygon or a component force expressed in millivolts; thus each segment is converted to millivolts times sec/100.

Since the difference in time of onset for endocardial excitation for different muscle regions is not great, it is practical to begin at 00 sec for the onset of action for a force. Thus the side of the polygon from 04 sec to 05 sec has an average duration of 045 sec. The length of the side depicts the magnitude of the force, e.g., 3 mv. In such a case $3 \text{ mv} \times 045 \text{ sec} = 135 \times \text{sec}/100$.

This value represents the force in units of time manifested by this one side of the polygon. A similar procedure should be done for each successive side. The sum expressed in $\text{mv} \times \text{sec}/100$ is an index of the voltage in units of time created by the left ventricle for one cardiac cycle.

In order to be comparable from one subject to the next the index of maximum potential and the index of potential seconds must be converted to terms of electrical moment by multiplying by the factor $R^2/3L$. This factor is the reciprocal of the sensitivity the transverse lead used as a

basis for standardization. The index of maximum potential and index of potential seconds are determined per cardiac cycle, to be expressed per minute they must be multiplied by the pulse rate (Fig 89)

THE NORMAL VECTORCARDIOGRAM

The normal vectorcardiogram depicts a QRS loop to the right at birth. It is of small magnitude in terms of electrical moment (Fig 90). There is a gradual transition to the adult type vectorcardiogram. In the child the values obtained from the VCG (index of maximum potential and index of potential seconds) remains small (Fig 91). The values increase with increasing body size until early adult life. From this peak there appears to be a decline in the size of the VCG. The average forty year old American male has a decrease in index of maximum potential. The significance of this is not clear. Theoretically it could represent changes in density of charge across the wave front and is related to cell function or even a decrease in cardiac output. Common types of normal vectorcardiograms are illustrated for the reader to compare with the routine ECG (Figs 92, 93, 94 and 95).

The VCG in athletic individuals is usually much larger with a large index of maximum potential and a large index of potential seconds per cardiac cycle. These individuals commonly have slow pulse rates with large stroke volumes. The surface area of the ventricles is greater increasing the size of the excitation wave front manifested by an increased index of maximum potential. The ST interval is often displaced and the T loops are increased in magnitude. The increase in T magnitude is evidence of physiological enlargement rather than pathological change.

Characteristically the QRS loop of the athlete is opened consistent with cardiac dilation. The terminal portion of the loop is often complicated indicating a complicated wave front at the base of the dilated ventricle (Figs 96 and 97).

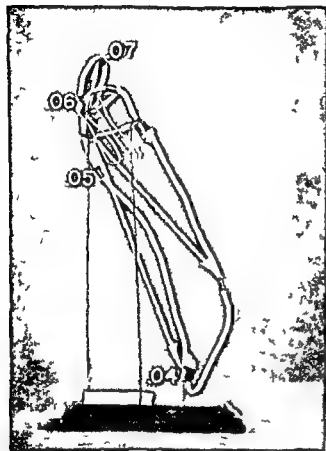


Figure 89 In the above example the QRS duration is .09 sec (the .08 time marker is behind the T loop). The long left ventricular vector occurs at .04 sec. The length of the pathway from .04 to .09 sec is the index of maximum potential. In this case the length is 31.5 cm or 299.25 mv. (1 cm equals 9.5 mv in terms of electrical moment). The lengths for each time interval for calculation of potential seconds are as follows:

04 - 05 sec	= 16 cm	= 152 mv	= 684 mv sec/100
05 - 06 sec	= 6 cm	= 57 mv	= 313.5 mv sec/100
06 - 07 sec	= 4.5 cm	= 42.8 mv	= 278.2 mv sec/100
07 - 08 sec	= 4 cm	= 38 mv	= 305.0 mv sec/100
08 - 09 sec	= 1 cm	= 9.5 mv	= 80.8 mv sec/100

1661.5 mv sec/100

or 1.66 volts \times sec/100 equals the potential seconds per QRS cycle. At a cardiac rate of 80 per minute the potential seconds per minutes would be 1.66 volts \times sec/100 \times 80 or 132.8 volts \times sec/100 or 1.33 volt sec per minute.

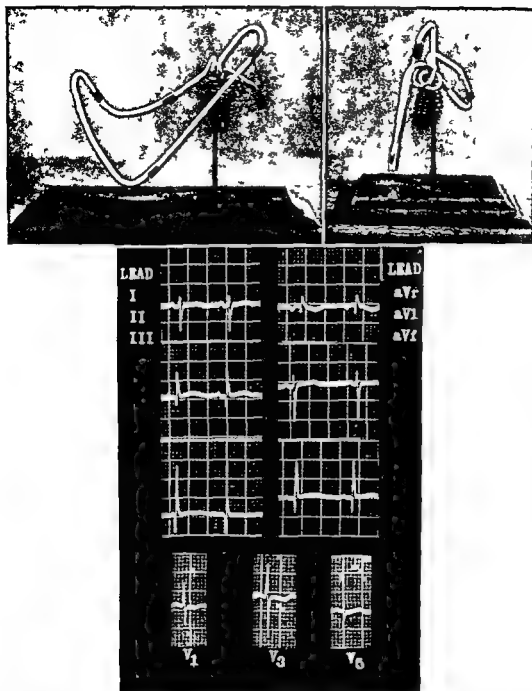


Figure 90 A vectorcardiogram of a seven day old infant. Note the rightward direction of the QRS loop and the mean QRS axis on the ECG of $+120^\circ$. The initial R wave in Lead I is depicted by the initial leftward course of the QRS loop. The terminal S wave depicts the rightward course of the loop. Try to reconstruct the character of the QRS complex in each of the other six limb leads by projecting the loop on the correct lead axis. The QRS duration is only .05 sec. This model is the only one not corrected for electrical moment as it was recorded by a different technique in 1951.

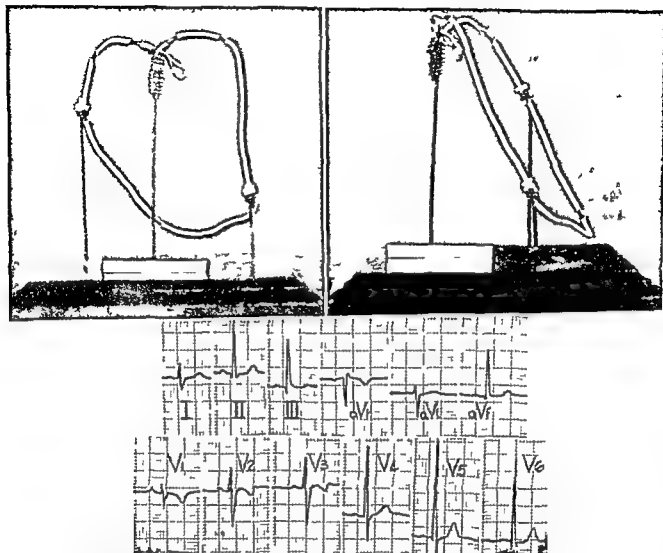


Figure 91 The VCG of an eight year old girl. The QRS loop is small in terms of electrical moment. This is an excellent example of small QRS amplitude when expressed in terms of electrode placement. The QRS amplitude on the routine ECG is not expressed in terms of electrode placement and the QRS amplitude therefore bears no relation to the true size of the small heart of this 56 pound child. The small P loop is directed posteriorly and obscured by the larger T loop. The T loop is directed posteriorly on the VCG and is manifested by an inverted T wave in V_1 , V_2 and V_3 . This is the juvenile pattern. The spatial VCG suggests that the apex of the heart may be pointed backward in such examples.

The vectorcardiogram provides a clear demonstration of the influence of respiration. The expiratory VCG consistently is the largest. At the onset of inspiration there is marked decrease in size of the last half of the QRS loop representing the left ventricle and slight increase in the anterior segment. When the breath is held in deep inspiration the left ventricular component momentarily increases with the surge of blood

from the lungs into the left ventricle but there is an overall decrease in the loop consistent with decreased stroke volume (this is most marked in complete valsalva). On expiration the VCG is momentarily larger than during quiet breathing. The temporal changes in the QRS cycle do not permit the simple explanation of change in cardiac position due to excursion of the diaphragm (Fig 98).

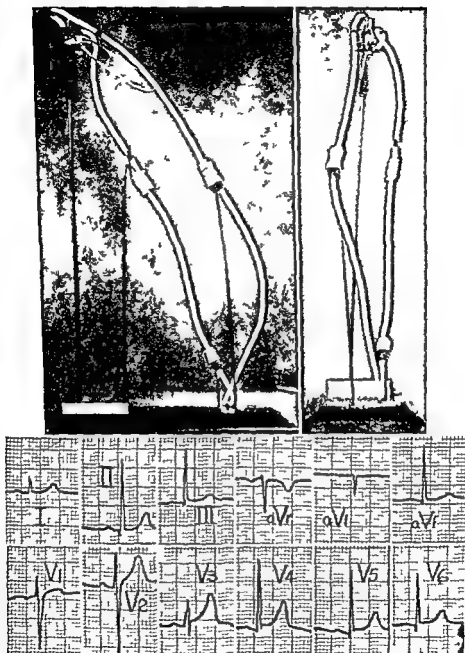


Figure 92 The VCG of an eighteen year old healthy male. The initial upward course of the QRS loop creates an initial R wave in aVR. The QRS loop remains open indicating a force is still active at the completion of excitation. This is the same as the elevation of the ST segment seen in Leads I, II, III, aVF and V2 through V6. The QRS loop continues directly into the T loop. The P loop is very small and directed nearly straight to the left.

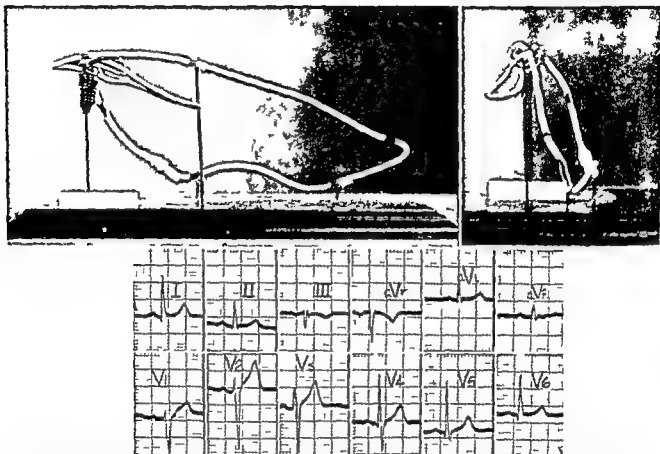


Figure 93 The VCG of a forty three year old man. The QRS loop is intermediate in location. The initial rightward course of the QRS loop corresponds to the initial Q wave in Lead I and the initial R wave in Lead III. The subsequent leftward QRS pathway writes an R wave in Lead I and an S wave in Lead III. When the QRS pathway descends it enters the positive zone of Lead III creating an R wave. The terminal QRS pathway is directed upward to such a degree that a terminal S wave is inscribed in Lead III.

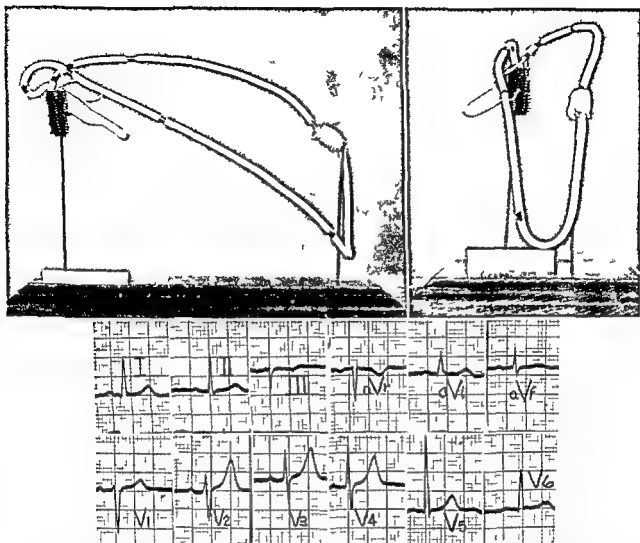


Figure 84 An intermediate VCG in a normal forty year-old man. The QRS pathway is first directed to the right and downward (Q wave in Lead I and embryonic R wave in Lead III). As the pathway is directed upward a tiny S wave is inscribed in Lead III. The QRS pathway continues to the left and the inferior limb of the main QRS loop. It invades only a little way into the positive zone of Lead III creating an R wave of very low amplitude. The terminal QRS pathway is well within the negative zone of Lead III creating a more prominent S wave. The chief difference in this VCG from Fig. 93 is that the main terminal QRS pathway is above the main initial large QRS pathway; thus there is no R wave following the main S wave in Lead III. The initial R wave in the V leads becomes progressively larger as the V lead axis becomes more leftward. Compare the configuration of Lead I and V6.

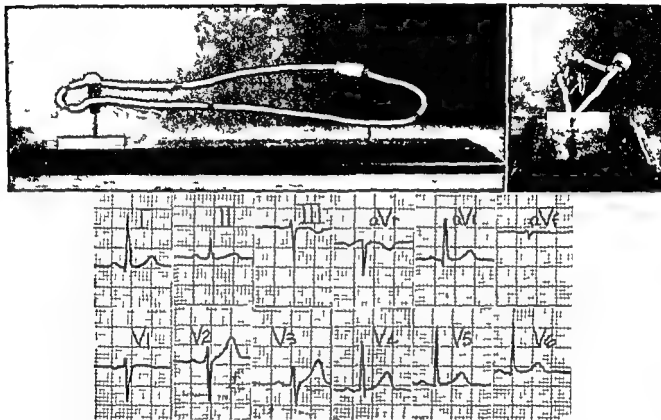


Figure 95 A VCG with horizontal orientation. The QRS loop is narrow and nearly perpendicular to V_1 . This accounts for the low amplitude and diphasic character of the QRS loop in that lead. The loop is also narrow in the sagittal view corresponding to the low QRS amplitude in V_2 . The area of such long narrow loops is quite small. Calculations based on the enclosed area are meaningless. In such examples only the correct principles of the polygon sides (length of QRS pathway) can be expected to reflect the magnitude of electrical activity. The P loop as usual is smaller than the T loop.

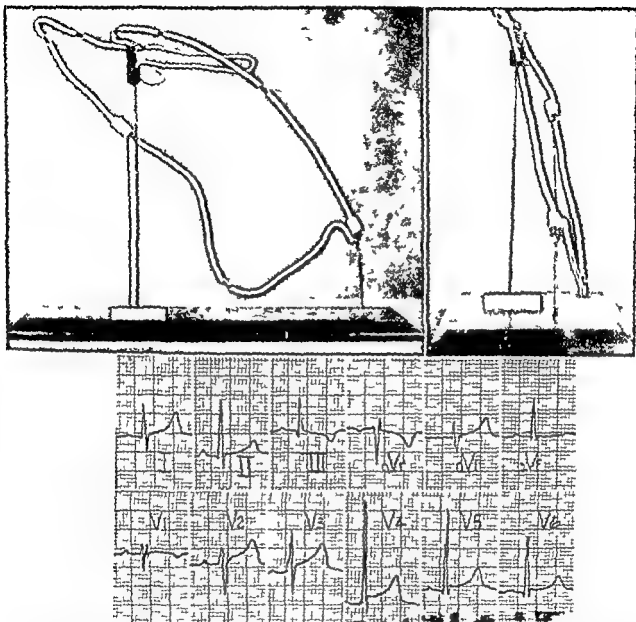


Figure 96 The VCG of a twenty seven year old Olympic athlete. The QRS configuration of Lead I can readily be extracted from the frontal plane. Regard the 5 cm standardization to appreciate the tremendous size of the VCG. The marked increase in QRS amplitude is lost in the routine ECG. The subject has a resting pulse rate of 44 per min. with necessarily increased stroke volume. The ST interval creates an open QRS loop. The T loop is large and normally directed. The smaller P loop is clearly evident.

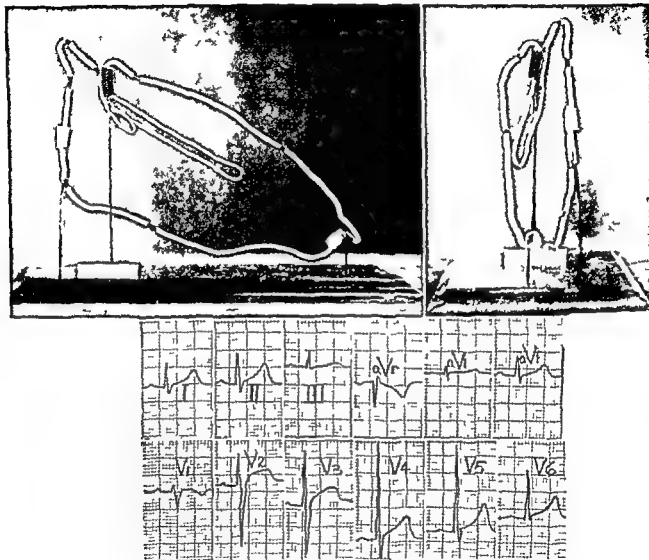


Figure 97 The VCG of a thirty seven year old Olympic athlete pulse 46 B P 92/52 The large QRS loop is clearly evident associated with a large T loop and smaller P loop The QRS loop is open

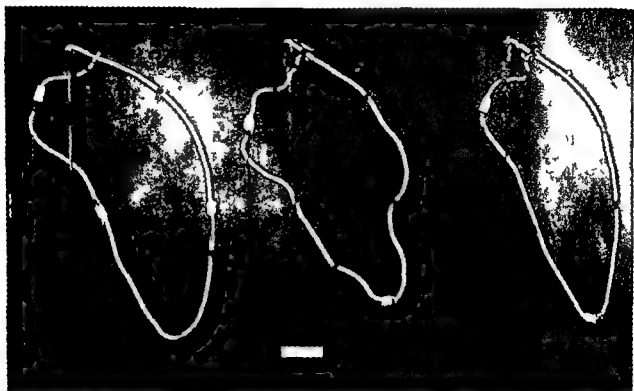


Figure 98 The VCG shows marked changes during the respiratory cycle. The above models are all of the same individual. On the left the subject is in expiration. The center model represents the early phase of inspiration. The model on the right shows the increase in size of the VCG that occurs while holding the breath in full inspiration despite the fact the lungs are at maximum inflation. Note the absence of rightward shift of the VCG and the selective changes in the last half of QRS pattern on early inspiration.

IX

Cardiac Enlargement

ATRIAL ENLARGEMENT

ATRIAL ENLARGEMENT is manifested by an increase in duration or amplitude of the P wave. It is not possible to detect which atrial chamber is enlarged from the routine electrocardiogram. However, there is a tendency for a more leftward P loop or P axis with left atrial enlargement and a more vertical loop with right atrial enlargement (Figs 99 and 100).

LEFT VENTRICULAR ENLARGEMENT

As the left ventricle enlarges the length of time required for completion of a confluent wave front of excitation is increased. This is caused by the increase in endocardial area to be excited or directly related to ventricular dilatation. This factor can increase the time required

for excitation (increased QRS duration) without significant increase in wall thickness.

Dilatation of the left ventricle increases its surface area and consequently increases the size of the excitation wave front. This is manifested by increased QRS amplitude, increased index of maximum potential and increased potential seconds. The septum remains as a 60° arc of the ventricular cone. Elongation of the ventricle increases the septal area (in relation to the base). The increased septal component opens the spital QRS loop creating a prominent posterior component during the last half of the QRS cycle. The open loop is characteristic of ventricular dilatation and elongation. Opening of the loop cannot be determined on the routine ECG. It is related to the amplitude and difference in time between the S wave in V2 and the R wave in V6.

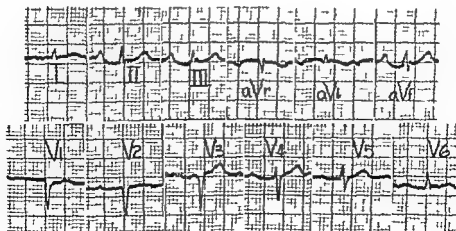


Figure 99 The ECG of advanced pulmonary emphysema. The PR interval is prolonged to .24 sec. The mean P vector is $+90^\circ$ with an isoelectric P interval in Lead I. The mean QRS axis is $+60^\circ$. Note the low amplitude of the QRS complexes throughout the record. The distinctive features are the comparatively large P waves with vertical P axis and the very small QRS amplitude

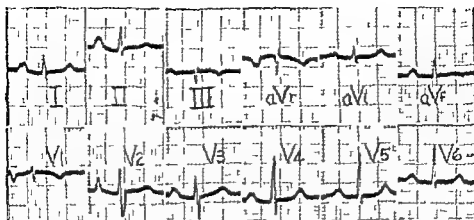


Figure 100 P mitral This is an example of mitral stenosis and insufficiency PR interval 26 sec increased P amplitude P axis +45 comparatively small QRS amplitude mean QRS axis +15 The distinctive feature is the large P waves in Lead I and II

Amplitude is often misleading when measured by the routine ECG due to electrode distance and lead length. Whenever the sum of the amplitude of the QRS complexes in Leads V6 V2 and aVf exceeds 5 millivolts the question of dilatation should be considered. A prominent S wave in V2 is a frequent finding in left ventricular hypertrophy.

Generalized dilatation may be manifested by increased voltage. The increased magnitude seen in the presence of a normally slow heart is such an example. There is an increase in amplitude in both the R wave and S wave in V2. In isolated left ventricular hypertrophy the R wave is normal or diminished in amplitude whereas the S wave is increased in size. The increase in voltage caused by uncomplicated left ventricular hypertrophy affects chiefly the last half of the excitation cycle or that portion dominated by the left ventricle.

The value of converting measurements to terms of electrical moment is evident in comparing such a vectorcardiogram in a child or normal adult and in the presence of hypertrophy (Fig 101). It is known that the child's heart is small and the relative size difference can be appreciated in this manner.

Increase in wall thickness increases the QRS duration and the potential seconds. The QRS duration becomes more significant as an index

of wall thickness in the absence of dilatation (normal QRS amplitude).

Ventricular recovery is altered in the presence of left ventricular hypertrophy. With increased wall thickness the time required for transmission of the wave front of recovery to the surface is increased. The early endocardial recovery at the apex creates a wave front before arrival at the epicardial surface. This wave front directs forces towards the recovered endocardial tissues or towards the base. The apical endocardial recovery creates an ST segment depression in those leads with an upright QRS complex. The degree of depression depends upon the degree of wall thickness.

It is interesting to note that physiological enlargement as seen in the athlete is often accompanied by ST segment changes. These however are due to a vector directed parallel to the QRS vector and are associated with early recovery of the right ventricle and apical regions throughout the thin wall.

With increased wall thickness the conical wave front of recovery develops steeper walls or a cylindrical shape. This diminishes the magnitude of the T loop and T waves. Progressive wall thickening results in an everted conical wave front throughout the endocardial shell. At this point the T vector is directed opposite the QRS vector or creates a wide spatial QRS-T

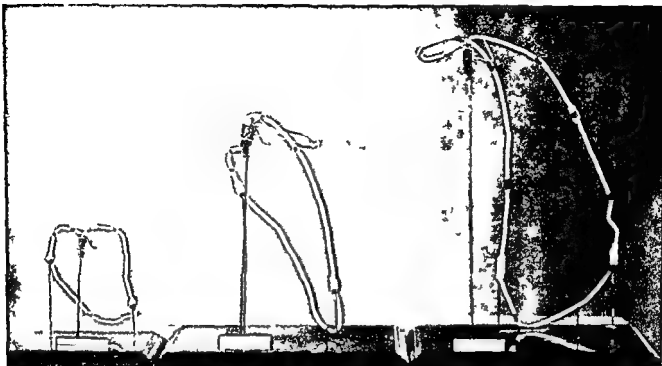


Figure 101 Three VCG models constructed to the same scale (see standardization bar). The left model is a fifty six pound child. The center model is a normal twenty two year old 155 pound male. The model on the right is that of a twenty-one year old female at 120 pounds with congenital aortic stenosis and marked left ventricular enlargement. The difference in cardiac size is unquestioned clinically and the difference in the VCG model size is obvious.

angle the T loop is opposite the QRS loop the QRS complexes are upright with inverted T waves. The inverted T wave becomes progressively larger after this stage.

The increased time required for excitation and recovery causes an increase in the QT interval. Physiological enlargement which is chiefly dilatation is associated with a large normally directed T loop (Fig 102).

Left ventricular hypertrophy may be present with vertical intermediate or left electrical axis (Figs 103 and 104). Marked left axis is more apt to occur with elongation of the ventricle in association with other physical factors. A high incidence of angina, previous infarction and conduction defects occurs in subjects with an electrical axis further left than -30° .

Left ventricular hypertrophy causes *

1) Increased QRS duration (increased potential seconds)

2) Increased QRS amplitude (increased index of maximum potential)

3) An ST vector directed away from the QRS vector. Physiological dilatation creates an ST vector parallel to the QRS and T vectors.

4) Decreased T vector magnitude. Physiological dilatation increases T vector magnitude.

5) Isoelectric T waves

6) Inverted T waves with a wide QRS T angle. Physiological dilatation is associated with a normal directed T wave.

7) Increased QT interval

RIGHT VENTRICULAR ENLARGEMENT

Enlargement of the right ventricle creates particularly complicated electrocardiographic

length) the difference in opinion amongst authorities regarding what is left ventricular hypertrophy by the ECG. This is hardly surprising since the important factors of magnitude are not considered (electrode distance and lead

rate). The difference in opinion in determination and use of the so called intrascenic deflection the failure to use the sides of the polygon principle to evaluate magnitude and the absence of any correlation of QRS magnitude to cardiac

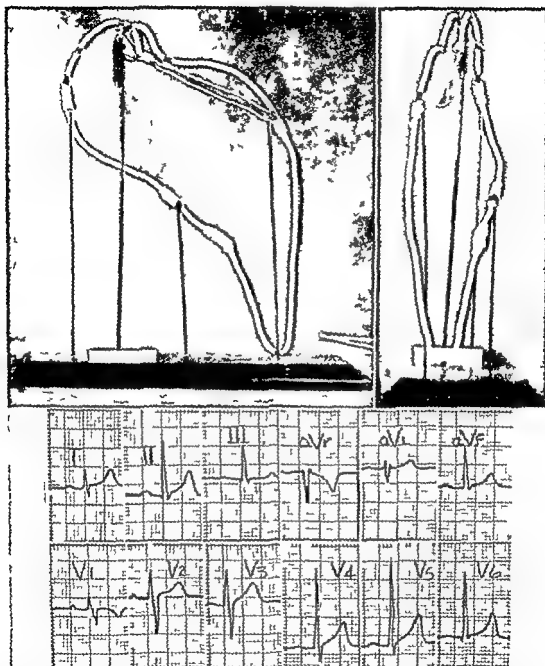


Figure 102 The VCG of an Olympic athlete. The large normally directed T loop is characteristic of physiological dilatation. The ST interval is manifested by a normally directed ST force. There is an overall enlargement of the entire QRS loop as one would expect from generalized dilatation. Thus the R wave in V2 is large compared to the S wave. Contrast this to the R/S ratio in hypertrophy due to selective increased work of the left ventricle in the subsequent figures.

manifestations. In addition to a simple consideration of chamber size and wall thickness there is the problem of change in position of the left ventricle secondary to right ventricular dilatation.

Simple uncomplicated right ventricular dilatation pushes the left ventricle posteriorly (Fig 105). This simple maneuver causes the terminal events of left ventricular excitation to describe a spatial pathway to the right and posterior. It requires only a small backward shift of the apex to accomplish this effect. The left ventricle then creates a larger S wave in

Lead I with a vertical or rightward mean QRS axis. There is an S wave in Lead V2. The terminal rightward pathway may describe an R wave at V1.

As long as the left ventricle is the last to complete excitation the S wave in Lead I cannot be ascribed to the right ventricle. When the right ventricle dominates the terminal excitation there will be little if any S wave in Lead V2.

Dilatation of the right ventricle causes prolongation of the time required to form a confluent wave front of excitation. This prolongation in

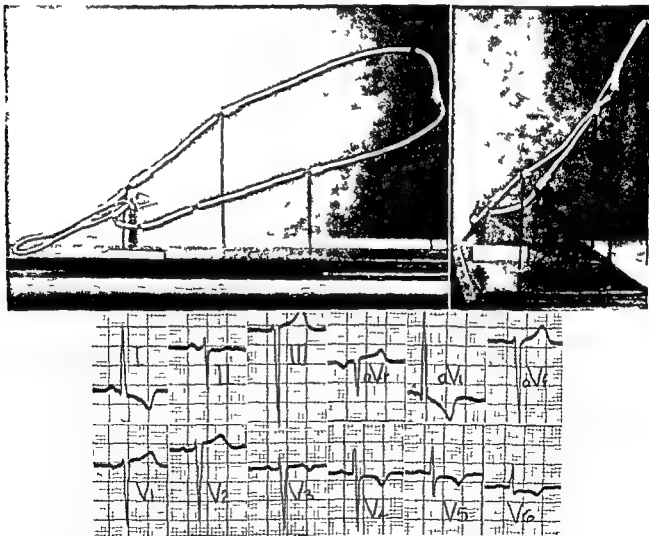


Figure 103 The VCG of left ventricular hypertrophy in hypertension. The QRS amplitude is markedly increased. The QRS loop is directed backward. The large T loop is directed opposite the QRS loop. Note the ST interval is directed parallel to the T loop. The V leads are recorded at half standard.

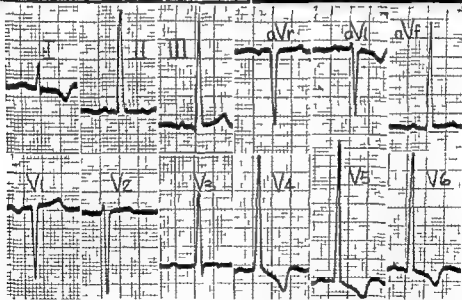
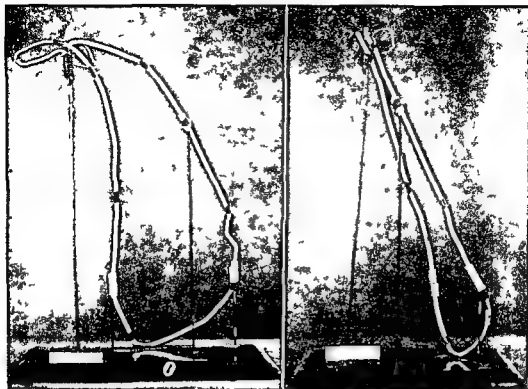


Figure 104 The VCG of twenty one year old case of aortic stenosis. The QRS loop is vertical but again chiefly posterior in its orientation. The T loop is comparatively smaller than the previous example and is directed opposite the QRS loop. The P loop is directed backward. Leads V2 and V3 are recorded at half standard.

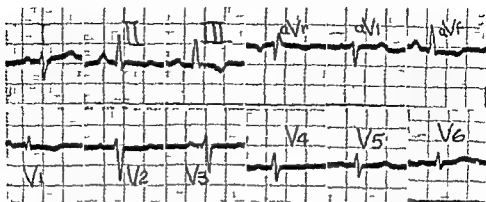


Figure 105 Right ventricular dilatation with a mean QRS axis of $+105^\circ$. The terminal S wave in V2 and Lead I represent the terminal forces of left ventricle. An S wave in Lead I due to the right ventricle would not have an S wave in V2.

time of excitation may allow the right ventricle to create a large force during the period normally occupied by left ventricular excitation alone. This opposite force diminishes the QRS amplitude and the index of maximum potential.

The thicker base may be sufficiently delayed in activation to create a late R wave from the right ventricle alone. In this case right ventricular hypertrophy resembles the electrocardiographic pattern of right bundle branch block (Fig 106). It differs from bundle branch block in that its index of maximum potential is usually less.

Right ventricular dilatation may cause

- 1) Increased QRS duration
- 2) Decreased index of maximum potential

3) Right axis shift of mean QRS axis manifested by a prominent S wave in Leads I and

V6 terminal R wave in V1 and terminal R in Lead III.

4) Delay in activation of the base with complicated conduction resembling bundle branch block.

When right ventricular dilatation is associated with increased wall thickness there is further prolongation in its activation. The prolonged period of activation may maintain a larger wave front in the right ventricle than in the left. This creates right ventricular dominance of excitation. As left ventricular activation is completed the right ventricular force directs the spatial pathway anterior. If right ventricular dominance is maintained the entire pathway is displaced anterior to the point of origin.

In this instance the rightward axis with terminal S wave in Leads I and V6 is due to the



Figure 106 An ECG of an adult with anomalous pulmonary venous drainage into the right atrium. The apparent right bundle branch block may disappear following surgical correction of the cause for right ventricular hypertrophy in such cases.

right ventricle. The QRS complex in V2 is chiefly positive with little or no S wave. These findings are most often seen in congenital lesions placing the increased work chiefly on the right ventricle — e.g., atrial septal defect and pulmonary stenosis. Since the right heart is larger at birth, the disproportion persists and the left ventricle never has an opportunity to exceed the size of the right ventricle.

Mitral stenosis with advanced right ventricular hypertrophy can develop an anterior loop with a completely positive QRS at V2. Left ventricular atrophy may contribute to the imbalance. When the left ventricle remains of normal thickness or if other complications are present, the S wave in V2 will be more prominent. Decrease in the index of maximum potential is almost constant in the presence of

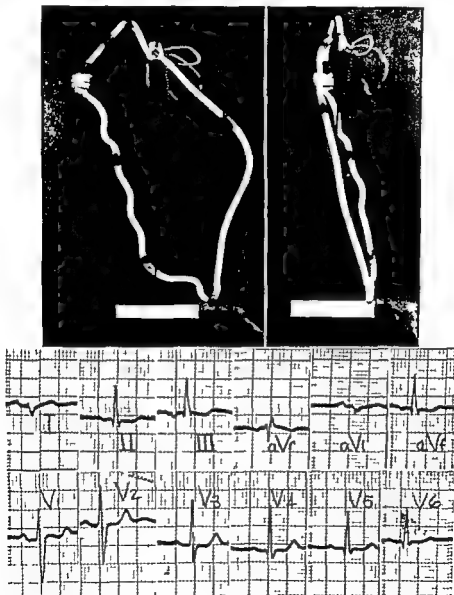


Figure 107. Mitral Stenosis. Note the vertical QRS loop, small QRS amplitude and prominent early anterior pathway of the QRS loop (prominent R wave in V2). The P loop is the most leftward loop (mean P axis -15°). The leftward P loop (left atrium) and vertical QRS axis suggests mitral stenosis.

significant mitral stenosis. The possibility of myocardial damage due to rheumatic activity could be an associated factor (Figs 107 and 108). As right ventricular hypertrophy ensues be-

coming the dominant ventricle with a thick wall it affects ventricular recovery much as does left ventricular hypertrophy. The T vector rotates away from the QRS forces widening the mean spatial QRS-T angle.

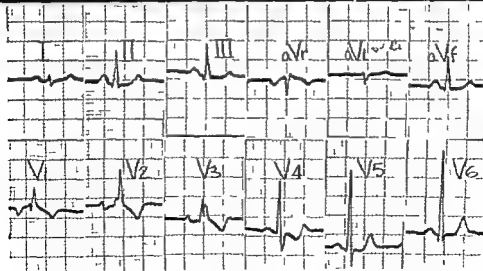
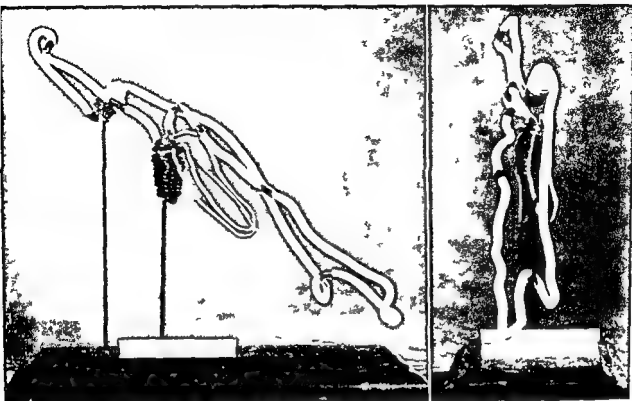


Figure 108 Advanced Mitral Stenosis. The small QRS loop is directed anteriorly (R wave in V2). The P loop is approximately as large as the T loop.

X

Conduction Defects

LEFT BUNDLE BRANCH BLOCK

THE SIMPLEST FORM of left bundle branch block is a single localized interruption of the left bundle with the rest of the endocardial conduction system intact. Due to the block in the left bundle the excitation impulse passes through the right bundle from the AV node. This causes excitation to begin at the right endocardial region at the septal area (Fig. 109).

Activation of the right septal surface creates a force directed posterior and slightly to the left (0.05 sec). When the right ventricular cone of activation is complete (0.12 sec) the resultant force is more leftward and commonly posterior. Excitation spreads outward from all regions. The free wall of the right ventricle gradually completes activation causing the instantaneous vector to be directed even more posteriorly. The excitation wave reaches the endocardial surface of the left ventricle at its thinnest region at 0.14 sec. As soon as the impulse reaches the left ventricle rapid endocardial excitation occurs. As the left ventricular cone becomes confluent the instantaneous vector moves leftward retaining its posterior direction. By 0.16 sec the wave front is the 300 arc left ventricular cone. The magnitude of the vector gradually diminishes to zero as excitation proceeds to completion.

The order of excitation in left bundle branch block commonly creates an early posterior pathway. The normal loop creates an early anterior pathway. Thus the loops of left bundle branch block are usually just opposite in direction to the normal loop.

The duration of excitation is prolonged to

0.12 sec. The free wall of the left ventricle normally requires 0.08 sec for excitation. The free wall does not receive activation until the impulse crosses the septum which requires 0.04 sec in the presence of complete block. Thus the delay in septal activation is responsible for the prolonged excitation.

Recovery begins at the right ventricular region. The formation of an endocardial wave front near the apex creates a force directed toward the base of the right ventricle. As recovery ensues the free wall of the right ventricle completes activation. The septum undergoes recovery from both sides, thus its force is small or non-existent. The left ventricular free wall recovers from the endocardial region. The resultant vector of the wave front is directed toward the septum and left ventricular base (Fig. 110). Recovery of the right ventricle prevents the QRS loop from closing and subsequent recovery creates a spiral pathway opposite the pathway of excitation.

If one projects the instantaneous vectors of the spiral pathway created by left bundle branch block on the lead coordinates the pattern of left bundle branch block is created. Since the QRS loop is directed posterior the QRS complexes in V1 and V2 are commonly negative deflections and the leftward course creates a positive QRS complex in Leads I and V6. The onset of right ventricular recovery creates a force opposite the QRS loop or an ST segment depression in those leads with an upright QRS complex. The T loop is opposite the QRS loop and creates a negative T wave in leads with an upright QRS complex (Fig. 111).

The characteristics of left bundle branch

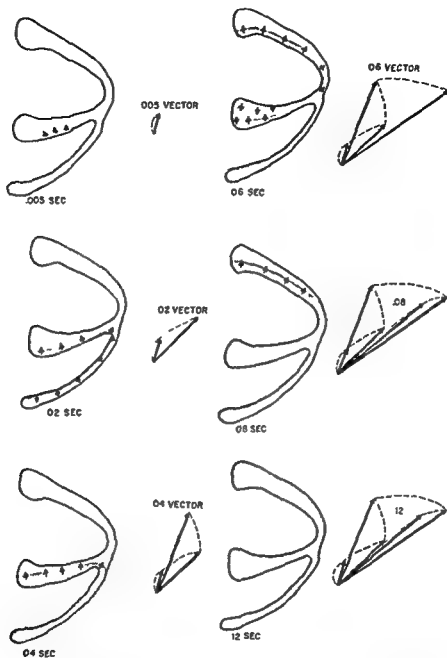


Figure 109 Order of excitation in left bundle branch block

block noted on the routine ECG are

- 1) QRS duration of 0.12 sec or more
- 2) Upright QRS complexes in Lead I and V6
- 3) ST segment depression in Lead I and V6 or an ST vector opposite the mean QRS vector
- 4) Inverted T waves in leads with upright QRS complexes i.e. a mean T vector directed opposite the mean QRS vector

In the event there is loss or absence of additional specialized conduction tissue in the left

ventricle activation through the left ventricle will be slowed. In its greatest extreme the entire left ventricle could require activation by slow muscle spread. This would greatly prolong the QRS duration beyond 0.12 sec.

The clinical significance of left bundle branch block depends upon its cause. It may be caused by arteriosclerotic heart disease or myocarditis. In very rare instances it is thought to be congenital in origin. Left bundle branch block

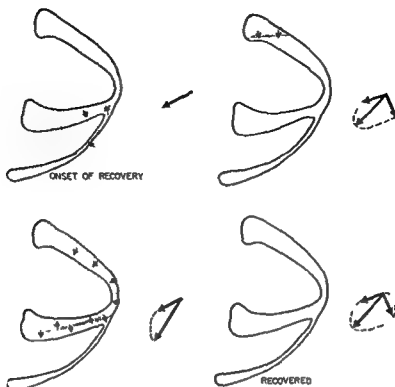


Figure 110 Order of recovery in left bundle branch block

without cardiac disease is almost nonexistent in children

RIGHT BUNDLE BRANCH BLOCK

Whenever septal activation is accomplished from only one side it will require twice the usual length of time. When the right bundle is blocked activation begins normally at the left endocardial surface and forms a normal left ventricular conical wave front. Right ventricular activation begins when this wave front crosses the septum. The thick base of the right ventricle is the last to be activated. The terminal force is to the right and anterior (Fig 112).

The initial forces of excitation are created by normal left ventricular activation and are manifested by an anterior pathway. There may be a very small short duration force directed to the right with the onset of septal activation. The remainder of the early pathway is leftward. The terminal pathway is rightward and anterior. The initial and terminal anterior direction of the forces creates an R R wave over the in

terior precordium at V1 and V2. The QRS complex at V1 and V2 may be entirely positive. The terminal rightward forces creates a terminal S wave in Lead I and usually a terminal R wave in Lead III (Figs 113, 114 and 115).

Recovery proceeds normally from apex to base creating a force directed toward the apex. Thus there is a normal upright T wave in Lead I. This has led to the observation that the T was upright in the lead with the terminal S wave (Lead I).

The characteristics of right bundle branch block are

- 1) QRS duration of 0.11 sec or more
- 2) Terminal S wave in Lead I and terminal R wave in V1 (a terminal right anterior vector)
- 3) There may be T wave changes secondary to the change in order of activation

Right bundle branch block seems to be more common than left bundle branch block in otherwise normal children. Its presence does not demand a bad prognosis. The problem is to detect underlying heart disease if any. Right ven

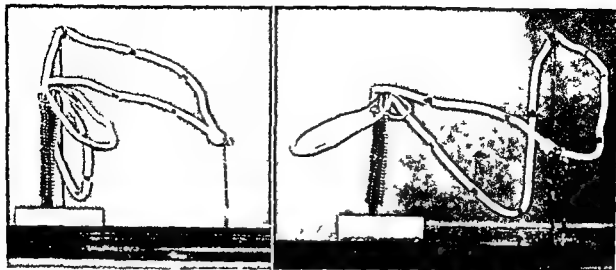


Figure 111 A thirty year old pilot with complete left bundle branch block of undetermined origin. Correlate the model with the earlier example of the order of excitation and recovery, then project model on lead coordinates to construct 12 lead electrocardiograms.

tricular dilatation and hypertrophy often create its similar or indistinguishable findings. The differentiation of right bundle branch block from right ventricular hypertrophy is not entirely satisfactory by current methods.

THE S S S PATTERN

A frequent finding in young people is the presence of a terminal S wave in Leads I, II, and III (Fig 116). This means the terminal force of excitation must be directed toward the right shoulder. In some instances this is associated with an rSr at V1 and in other instances with a terminal broad S wave at V1. This difference depends upon whether the terminal event is anterior or posterior.

For the most part this finding is without

clinical significance as long as the QRS duration is less than 12 sec. The S₁S₂S₃ pattern has been attributed to a number of causes. The most frequent cause cited is the late activation of muscle tissue near the tricuspid or pulmonary valves. It is interesting to note that the terminal site of activation in the dog's heart is the base of the septum. This in itself could produce an S₁S₂S₃ configuration.

Distinction between an S₁S₂S₃ pattern and right bundle branch block is not always possible as both are associated with a terminal rightward vector. However if congenital origin can be established distinction is not important. If late activation of muscle tissue near the tricuspid or pulmonary valve is the cause it is actually a delay in activation in the right ventricle.



SEPTAL ACTIVATION



SEPTUM



LEFT VENTRICULAR ACTIVATION



LEFT VENTRICLE



TERMINAL RIGHT VENTRICULAR
ACTIVATION



RIGHT VENTRICLE

Figure 112 Order of ventricular excitation in right bundle branch block

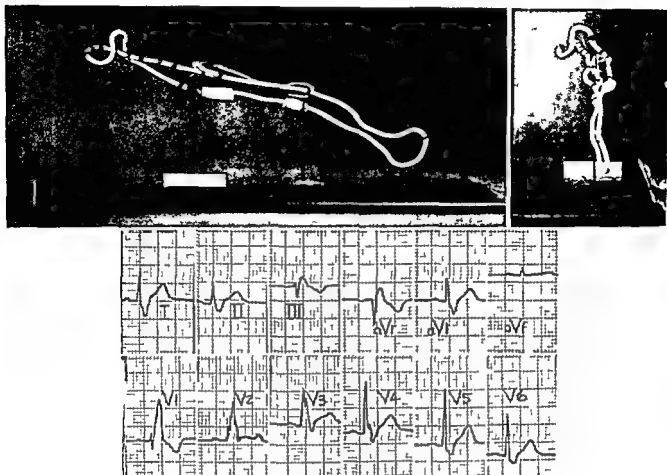


Figure 113 A twenty five year old pilot with a right bundle branch block. The close grouping of the time intervals depicts the conduction defect

INTRA VENTRICULAR CONDUCTION DEFECTS

A change in the order of excitation may occur at points other than the septum. The QRS configuration may be altered or the duration of excitation prolonged to nearly 12 sec. Occasionally these are called incomplete left or right bundle branch block. A better term is intraventricular conduction defect. The significance of this finding depends entirely upon its cause.

ACCELERATED CONDUCTION (Wolff Parkinson White Syndrome, False Bundle Branch Block, Pre Excitation)

Normally there is a delay between the completion of atrial excitation and the onset of ventricular excitation. This delay is manifested by

an isoelectric interval between the P wave and the QRS complex. In certain individuals ventricular excitation begins early or pre excitation occurs. Often a localized muscle area is stimulated early and the remainder of the muscle is excited in the usual manner. This shortens the PR interval and prolongs the QRS duration. The classic form is a PR interval of 0.10 sec or less and a QRS duration of 12 sec or more (Figs 117, 118, 119, 120 and 121).

Since the QRS duration is prolonged it resembles bundle branch block. The clue is the short PR interval. Pre excitation may occur in either the right or left ventricle and the QRS complex may resemble either right or left bundle branch block.

Two explanations have been advanced as cause for this finding. A bridge between the atria and the pre excited area called the Bundle

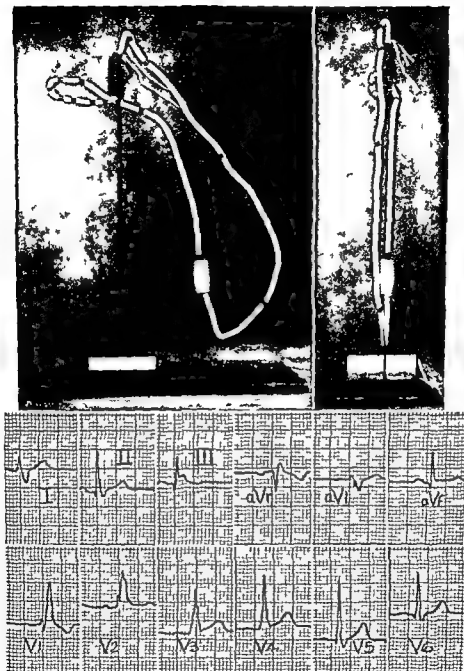


Figure 114 Right bundle branch block in an asymptomatic thirty four year old physician

of Kent was described as a means of early passage of the impulse around the AV node. Another explanation is to consider the AV conduction system as a central nervous system of the heart. An acceleration of conduction through one part of the node would stimulate its muscle receptor area early. This is called accelerated conduction. This has been demonstrated in the presence of nodal disease.

Regardless of the mechanism causing pre-excitation it is clear that the excitation impulse must first pass over the atria to the region of the AV node before pre-excitation can be set off by accelerated conduction or through the Bundle of Kent. If the atria requires 12 sec (normal value is 0.6 to 1.2 sec) for activation the pre-excitation phase must begin at 12 sec or after a PR interval of 12 sec. In the event atrial excitation requires only 0.6 sec a PR interval of 10 sec gives a relatively normal period for delay between atrial and ventricular activation. The absence of a delay between completion of atrial excitation and the onset of

ventricular excitation should be used to determine the presence of accelerated conduction rather than fixed values for the PR interval.

The muscle area undergoing pre-excitation creates a small force of slowly changing magnitude. This creates a slow early QRS pathway and on the LCG graph a slow onset of the QRS complex. The wave created by early onset is called the "delta" wave.

The T wave may be either normally directed or altered secondary to the change in the order of excitation. The accelerated cycles may not be constant varying with respiration or alternating from cycle to cycle. Occasional normal records are recorded at intervals in the same subject.

Accelerated conduction is usually thought of as congenital in origin. However accelerated conduction has been demonstrated in proved nodal disease. In hospital patients and those seeing a physician 50 to 75% have recurrent paroxysmal tachycardia.

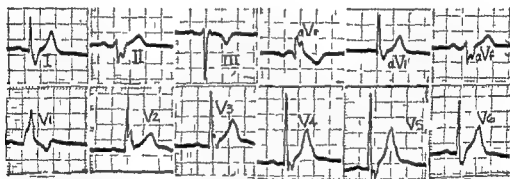


Figure 115 A thirty four year old pilot's electrocardiogram of complete right bundle branch block. An electrocardiogram during childhood showed an atrio-ventricular conduction defect which remained unchanged until age twenty-two. At age twenty-six an electrocardiogram showed complete right bundle branch block which has remained unchanged for eight years. Catheterization studies were normal as were the remaining parts of the examination. It is of interest that he had coccidioidomycosis.

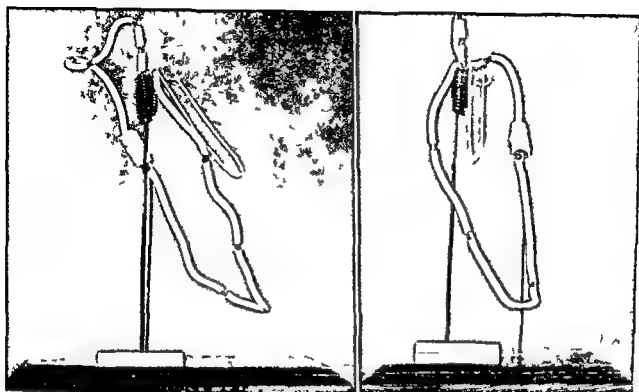


Figure 116 An S S S pattern in a normal adult. The terminal QRS pathway is directed toward the right shoulder creating an S wave in Leads I, II, and III. The close grouping of time intervals noted in right bundle branch block are not seen.



Figure 117 Accelerated nodal conduction in an asymptomatic twenty year old student. The short PR interval and prolonged QRS complex are characteristic. The prominent posterior course of the QRS pathway resembles left bundle branch block. The inverted T waves in Leads I, II and V6 are of no clinical significance.

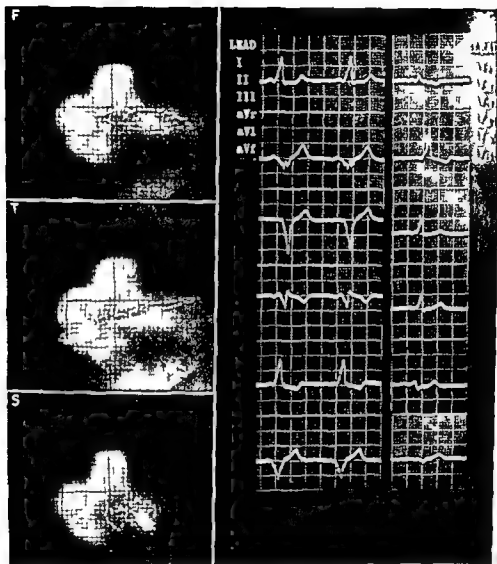


Figure 118 Accelerated nodal conduction in a forty year old woman. The slow conduction at the onset of the QRS is classical. The anterior orientation of the QRS pathway resembles right bundle branch block.

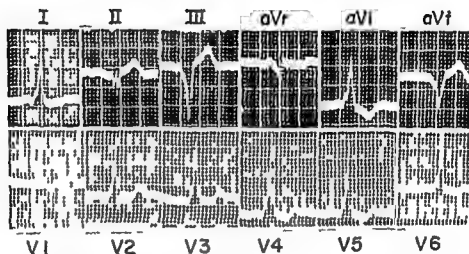


Figure 119 Accelerated nodal conduction in an eighteen year old boy found upon hospitalization for possible rheumatic fever. Subsequent investigation uncovered the same finding in records recorded during early childhood.



Figure 120 Upper left On admission the patient had atrial fibrillation Upper right Sinus tachycardia (150/min) The small P wave runs into the QRS complex in Leads II and III In V2 the sinus mechanism is readily apparent Lower left At normal rhythm the short PR interval of accelerated nodal conduction is seen The QRS duration is within normal limits Lower right The P wave occupies 12 sec causing the PR interval to be 12 sec There is no isoelectric interval between the P wave and the QRS complex

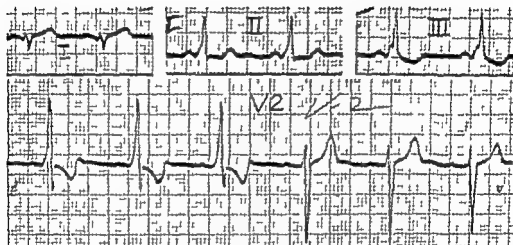


Figure 121 An example of accelerated nodal conduction that converts to normal conduction during recording of V2

XI

Pericarditis

ACUTE PERICARDITIS creates generalized epicardial injury. Injury current creates a force away from the zone of injury during the resting state and towards the zone of injury during the excited state. Thus the resultant vector of generalized epicardial injury is directed toward the base of the heart in the resting state and toward the apex in the excited state (Fig 122)

pression in leads with an electrode facing the base of the heart (aVr and VI)

The injury vector during the excited state prevents the forces of excitation from returning to zero value (base line or point of origin) consequently the QRS loop remains open. The spatial vector directed from the point of origin to the ST interval is the spatial injury vector

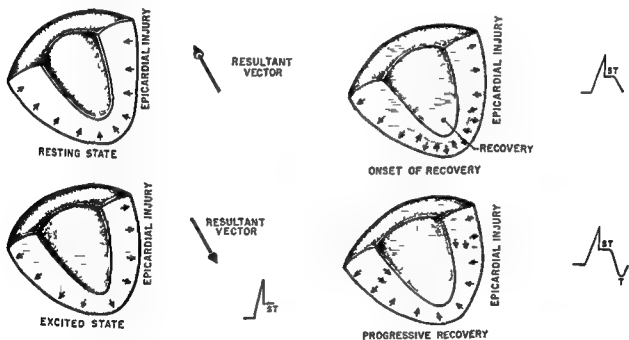


Figure 122

The injury vector merely displaces the base line during the resting state so that the injury force is apparent only during the excited state. This is manifested by ST segment displacement. The apical direction of the vector in the excited state causes ST segment elevation in those leads with positive electrodes facing the heart and de

pression in leads with an electrode facing the base of the heart (aVr and VI)

Recovery proceeds normally from the endocardial surface. Its onset causes a decrease in the resultant injury force or a decrease in ST segment elevation. When the wave front of recovery reaches the surface the injury force is

reversed towards the base of the heart (recovered tissue adjacent to injury has an injury force directed away from the area of injury). Thus the wave front of recovery forms a continuous shell with the epicardial region in the resting state. Effectively the wave front of recovery is an inverted cone.

The force of recovery is opposite the force of excitation and is manifested by an inverted T wave following an upright QRS complex or a T loop opposite the QRS loop. The T vector is

commonly small when the injury vector is large. As the injury vector diminishes in magnitude the T vector magnitude increases. The elevation of an ST segment in leads with an inverted wave has given rise to the rule that *abnormal ST segment deviations are present when the ST vector is directed opposite the T vector where normal ST segment displacement occurs with the ST vector and T vector directed in the same direction* (Fig 123). Serial records provide the best evaluation of the significance of ST segment displacement.

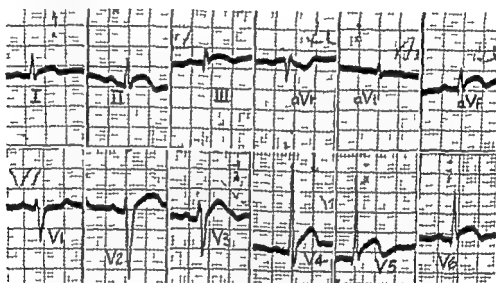


Figure 123 ST segment and T wave changes of acute pericarditis

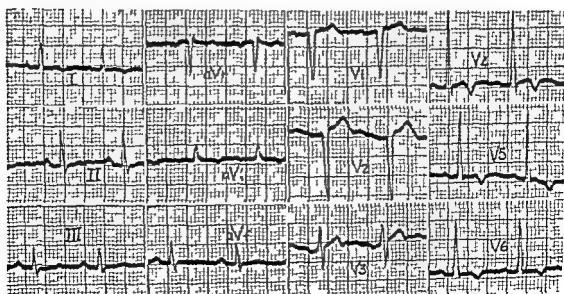


Figure 124 Chronic pericarditis with permanent T wave changes

As the area of injury subsides the magnitude of the injury vector diminishes. This results in closure of the QRS loop or return to normal of the ST segments. With complete healing of the injured area new cell membranes are formed and no constant current exists. T wave inversion may persist for an indefinite period of time following acute pericarditis and in some instances is permanent (Fig. 124). The scarred tissue over the myocardium creates local tension by pulling on the subjacent muscle tissue. Such mechanical action causes abnormal delay in epicardial recovery. Consequently the wave

front assumes the shape of an inverted cone with forces directed towards the base of the heart.

Pericarditis may be associated with decreased QRS amplitude. This may be caused by 1) actual muscle damage by inflammation or calcium deposits with diminished density of change on the wave front 2) increased heart rate in the febrile state or 3) cardiac compression diminishing cardiac volume and the size of the wave front either by effusion or constriction. Pericardial effusion occurs in the acute stage. Constriction occurs in the chronic form of pericarditis.

XII

Myocardial Infarction and Arteriosclerotic Heart Disease

MYOCARDIAL INFARCTION

INFARCTION OF THE MYOCARDIUM results from deficient blood supply to a muscle area. This may be due to increased need or decreased supply or a combination of the two. The most common cause of myocardial infarction is arteriosclerotic heart disease with coronary artery thrombosis. Coronary artery thrombosis is not entirely analogous to experimental ligation of the coronary artery because coronary thrombosis most often occurs in an individual with diseased arteries. The disease is not necessarily confined to the thrombosed vessel.

The myocardium receives its blood supply from the right and left coronary arteries. The arteries arise from the base of the aorta and extend over the ventricular surface. The right coronary artery and its ramifications extend over the posterior surface. The left coronary artery extends over the anterior and posterior ventricular surface. Ramifications of these arteries ex-

tend inward through the myocardial wall. The smaller vessels extending from the larger vessels create an interlacing network at the endocardial shell.

A number of conflicting viewpoints have been presented by various authorities to explain the ECG findings of infarction. To cite one major incompatibility consider the concept that in infarction involves chiefly the endocardial surface. If this is true the laws of injury current are incorrect. The forces of injury are directed toward the zone of injury during the excited state; therefore, a large endocardial infarction zone would create an ST force away from the zone of infarction (Fig 125).

The factual information based on empirical observations is as follows:

- 1) Infarction may occur without any significant changes in the routine ECG.
- 2) Infarction may be manifested only by T wave changes in the routine ECG (Fig 126).
- 3) The classical ECG pattern in infarction

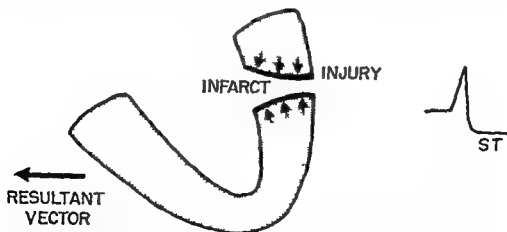


Figure 125 When endocardial infarction is larger than the epicardial region the ST vector is directed away from the site of injury rather than toward the infarction

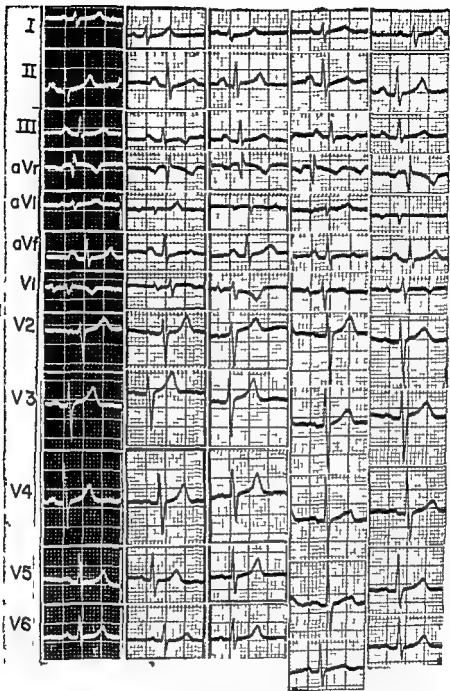


Figure 128 A) Base line ECG six months before illness in a fifty seven year old man B) Approximately 18 hours after minor chest discomfort The only significant change is the T wave in Lead III C) Three weeks later the T wave has gradually returned to its upright position There are minimal ST segment changes in Leads I II and V1 (not sufficient for a definitive diagnosis alone) D) Four weeks later following an episode of syncope while having his hair cut At four weeks and three days the electrocardiogram showed its maximum changes of minor T wave changes He had clinical infarction pain and labored findings following the episode of syncope E) Seven weeks from onset the ECG appeared almost identical to the base line record At eleven weeks he died suddenly without warning Autopsy findings demonstrated diffuse scarring of the anterior wall of the left ventricle with long standing disease of the anterior coronary artery The septum showed diffuse scarring of healed injury There was thrombosis of the right coronary artery

consists of an initial Q wave of 0.4 sec duration an elevated ST segment and an inverted T wave in those leads with a positive electrode facing the infarcted area. This is a changing pattern with gradual return toward normal and in some instances complete return to normal.

4) T wave changes persist in a portion of records and abnormal Q waves persist in others.

5) Infarction may present in initial finding of very large T waves that are upright rather than the classic inversion described above (Fig 127).

6) Infarction may decrease the QRS amplitude or the length of the QRS pathway.

7) Discrepancies in the location of the infarction at post mortem examination and the location indicated by the ECG can be attributed in part to eccentric zero center location and variation in lead lengths.

Bayley's* classic experiments on ligation of

Bayley R H and LaDue: Electrocardiographic changes of impending infarction and ischemia injury pattern produced in the dog by total and sub-total occlusion of a coronary artery. *Am Heart J* 28:94 1944.

the coronary artery have often been used as an explanation of the sequence of infarction. In considering the experimental findings it is well to remember that coronary artery disease was not present and presumably normal coronary flow occurred through the rest of the arterial tree.

The earliest change noted after ligation was inversion of the T wave. T wave changes were followed by ST segment elevation. If the vessel was tied a sufficient length of time a Q wave appeared.

It seems logical that if one considers the endocardial shell relatively deficient in blood supply it would be the first area to suffer from hypoxia under these circumstances. This would result in a larger upright T wave not an inversion. Injury should affect the endocardial shell first and thus would produce an ST segment depression. On the other hand if the epicardial shell is relatively deficient in blood supply the T wave and ST segment changes observed are compatible with the basic concepts of hypoxia and injury. The endocardial shell normally re-

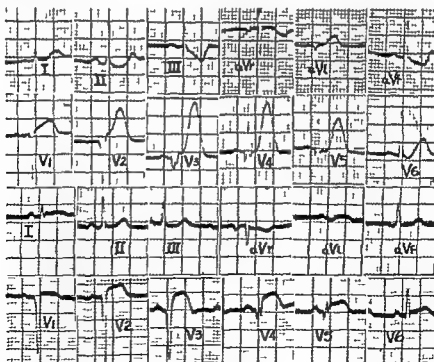
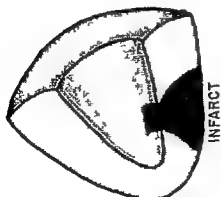
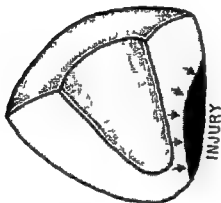
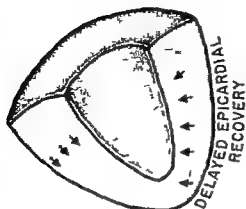


Figure 127 An example of large upright T waves at the onset of anterior infarction (above) and inverted precordial T waves in subsequent records (below)



**EXTENSION OF INFARCTION
TO ENDOCARDIAL SHELL**

Figure 128

ceives some blood supply from both major vessels. This probably accounts for the earlier changes in the epicardial region. If hypoxia persists, the entire thickness of the myocardial wall becomes infarcted. Involvement of the endocardial shell creates the abnormal Q wave (Fig 128).

A comparable situation occurs in the patient with localized coronary artery disease and thrombosis. Early T wave changes are followed by ST segment elevation and abnormal Q waves. The classic infarction may be defined in vector terminology as follows:

1) An initial QRS vector directed away from the area of infarction

2) An ST vector directed toward the area of infarction

3) A T vector directed away from the area of infarction

Using these rules, infarctions may be classified as to location. There are four principal locations: anterior, posterior, lateral, and diaphragmatic. Combinations of these areas may be involved (Figs 129, 130, and 131).

The initial forces of excitation are directed away from the infarcted area by the hole the infarction makes in the wave front. This will be the dominant effect of a transmural infarction until the conical wave fronts begin to change, i.e., the completion of right ventricular

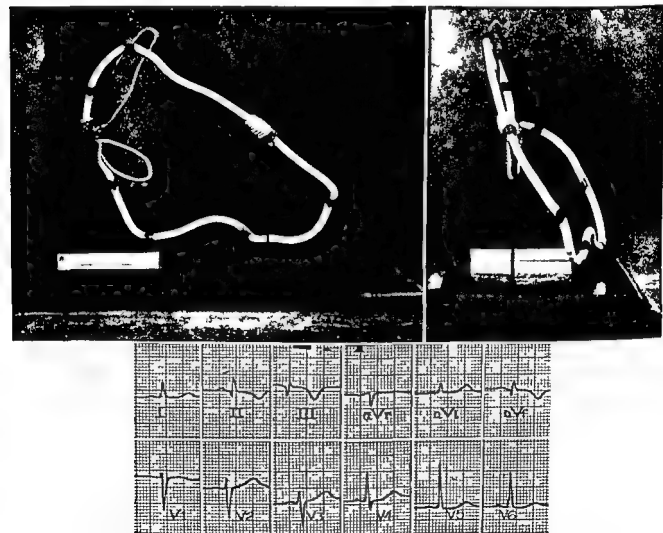


Figure 129 Infarction of the inferior surface of the left ventricle creates an upward initial course of the spatial QRS pathway. The T loop is directed away from the area of infarction. This is depicted by the Q wave and inverted T wave in Leads II, III, and aVF. The size of the QRS loop is small.

and septal activation. This period of time roughly corresponds to 0.4 sec. This has given rise to the clinical observation that infarction creates abnormal Q waves with a duration of 0.4 sec. Depending upon the location and size of the infarct, the remainder of the QRS pathway may be increased or decreased in size. After the wave front has passed the level of infarction, the QRS pathway assumes its normal course and magnitude.

Characteristic QRS changes are the most reliable hallmark of infarction. ST and T waves are important, but they are not always good indicators of the location of the infarction and may be due to many other causes.

The early large upright T waves may be explained on the basis of transmural hypoxia. In the event of shock or disease of other coronary vessels, the endocardial shell will not enjoy the additional blood supply from collateral cir-

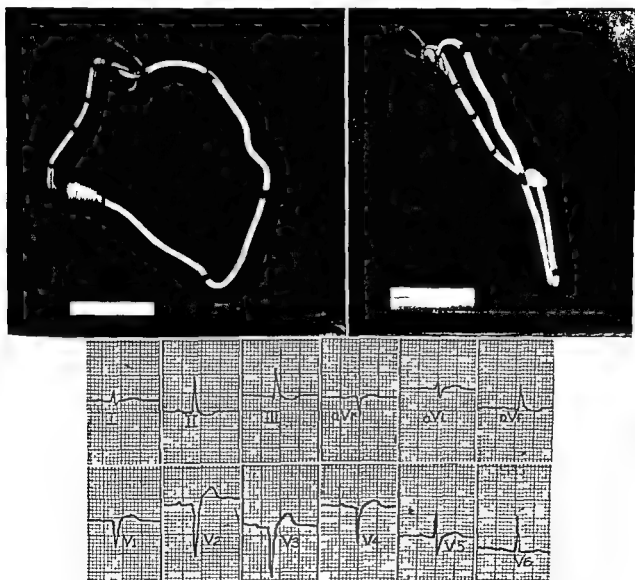


Figure 130 An example of anterior wall infarction. The QRS loop is directed entirely posterior. This is represented by QS deflections in I, II, III, aVR, aVL, and V1 through V4. The QRS loop is small. This individual's first evidence of infarction occurred eleven years ago.

ulation Occlusion of a vessel will cause in immediate hypoxia of the entire wall thickness followed quickly by injury and necrosis

Diffuse endocardial infarction is deserving of special comment Infarction of the entire endocardial shell will be apparent only as an ST injury force directed toward the base of the heart There is no hole in the wave front of excitation A complete infarction of the entire

endocardial shell without extension through the myocardial wall is rather rare (Fig 132)

The problem of endocardial infarction also concerns the depth of infarction Infarction extending only 3 millimeters into the wall will occupy a time interval of only 0.1 sec This time interval is so short it may not be recorded by the usual direct writing instrument

In order to alter the initial process of excita-

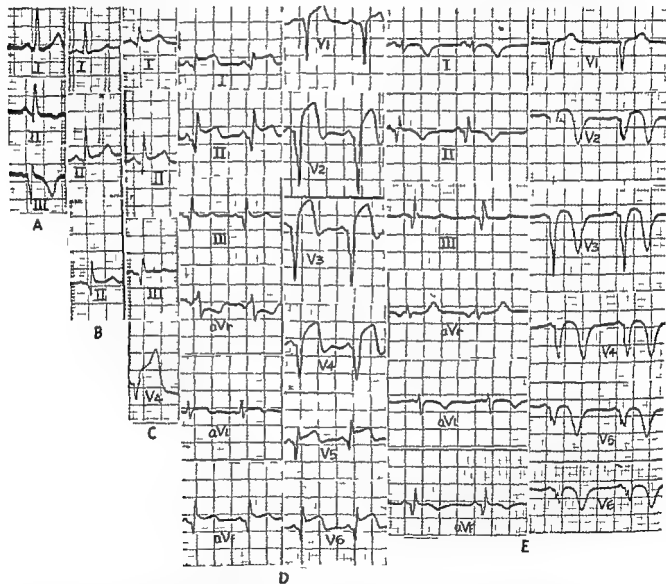


Figure 131 A) Electrocardiogram in 1947 five days after chest pain showing inferior wall infarction (Q, T) B) In 1952 the Q wave in Leads II and III persisted but were smaller C) One week later two hours after moving furniture Lead I and V4 show a new infarction D) The following morning a typical acute anterior wall infarction superimposed upon an old inferior wall infarction is evident E) Two weeks later the acute elevation of the ST segments has subsided and there are deep inverted T waves in Lead I and V2 through V6 The area involved is the anterior lateral surface of the left ventricle Note the loss of QRS amplitude in Lead I with the loss of the muscle area responsible for the leftward force of excitation

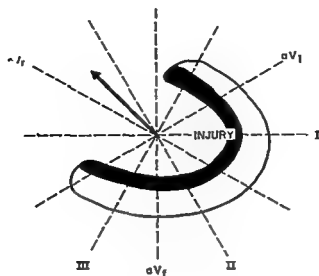


Figure 132

tion infarction must involve the endocardial shell. The duration of the influence of infarction depends upon the distance the infarction extends outward. The magnitude of the change in force depends upon the area of infarction. A small infarction creating a small hole in the wave front will not appreciably alter the direction of the resultant force of excitation.

The wave front of excitation for the left ventricle normally represents a cone at 0.2 sec. Its resultant force is perpendicular to the base of the cone or the base of the left ventricle. An infarction (hole) in the cone will tend to rotate the resultant. The new resultant force may be depicted as the resultant from the conical base

area and the infarction area (Fig. 133). The larger the hole, the greater is its effect on rotation of the resultant. One cannot expect the manifested forces during excitation to point directly away from the anatomic site of infarction unless the area of infarction is quite large.

Diffuse or patchy muscle necrosis creates no appreciable change in the direction of the forces. A localized area of necrosis within the wall thickness extending neither to the endocardial or epicardial surface will cause changes only midway in the excitation cycle. This is one cause for QRS notching.

Infarction involving only a portion of the outer myocardial shell will not be manifested

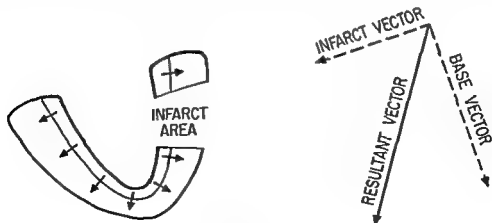


Figure 133

in the early portion of the excitation cycle. It will influence magnitude and direction of the later events of the cycle. Considering the above relationships, it is not surprising that infarctions may be manifested only by ST segment and T wave changes when studied by a routine ECG. While the ECG is a very valuable diagnostic adjunct in myocardial infarction, it does not replace the clinical diagnosis. Serial electrocardiograms demonstrating serial changes in conjunction with the clinical picture remains the most satisfactory approach at this time.

ARTERIOSCLEROTIC HEART DISEASE

Diffuse coronary artery disease causes changes in cellular metabolism. Diffuse fibrosis may occur. With loss of normally functioning cells, there is a decrease in ion density across the wave front of excitation. This can cause decreased QRS amplitude. The vectorcardiogram in terms of electrical moment clearly depicts the change (Fig. 129). A low QRS voltage after infarction favors generalized myocardial disease.

An occasional finding in arteriosclerotic heart disease is ST segment depression followed by an upright T wave. This has been explained as due to endocardial hypoxia. This finding is accentuated with exercise or during acute coronary pain. During an anginal attack there may be transitory T wave inversion. Acute ST segment depressions should always be respected as a possible indication of endocardial injury and impending infarction. Serial records should be obtained.

Arteriosclerotic heart disease may cause left axis and widening of the mean spatial QRS T angle. The T vector may be rotated more leftward than zero degrees. The QT interval may be prolonged.

At present the diagnosis of arteriosclerotic heart disease by the ECG in the absence of infarction or acute changes with angina is difficult and the tracing may appear normal. However, magnitude measurements have never been refined and measurements in terms of electrical moment should improve the diagnostic technique.

XIII

Drugs and Metabolism

DIGITALIS

DIGITALIS AFFECTS the basic electrical events of the cardiac muscle fiber. It alters the membrane action potential during the recovery phase. This phase is associated with an increased outward migration of potassium ions. Digitalis increases the permeability of the cell membrane to potassium ions, accelerating the outward potassium current. Clinically digitalis is given to increase the strength of muscular contraction. The refractory and contractile response of muscle fibers is closely related to intracellular potassium content. The effects on the recovery phase increase with increasing concentration of digitalis. At toxic levels the early period of the membrane action potential is altered, diminishing the initial spike. Since the early events of the membrane action potential are associated with an increased inward migration of sodium ions, the permeability of the membrane to sodium ions must be altered at these concentrations.

At therapeutic levels digitalis increases the speed of recovery of the ventricles. With more rapid spread of endocardial recovery, the wave front assumes the form of a large inverted cone. This produces a force toward the base of the heart during the ST interval (excited state). A normally directed resultant vector toward the apex is created when the wave front reaches the epicardial surface (Fig 134). This vector is diminished in size and duration due to the extensive early recovery of the endocardial shell. The larger the endocardial shell of recovery prior to recovery of the external surface, the larger will be the ST force and the smaller the

recovery (T) force. The enhancement of endocardial recovery produced by digitalis has three principal effects:

- 1) Shortening of the QT interval
- 2) Creation of a vector (ST) during the excited state in the opposite direction of the excitation (QRS) vector and opposite the normal recovery vector (T)
- 3) The larger the ST interval vector, the

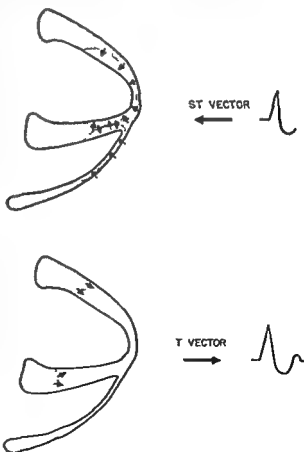


Figure 134

smaller the T vector. It is apparent that complete recovery throughout the endocardial shell will result in a short QT interval with a large ST force and little or no apparent T force (Fig 135).

The ECG findings of digitalis effect do not indicate intoxication nor can they be used as a reliable index for the proper dosage of digitalis.

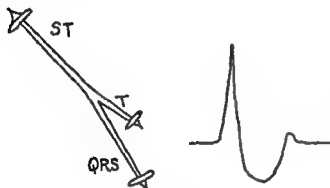


FIGURE 135

In toxic doses extra systoles may occur (Fig 136). Digitalis can cause multiple forms of arrhythmias. In toxic doses it may increase the PR interval but toxicity may occur before the PR interval is significantly prolonged. When the T vector has been previously altered due to other causes, e.g., left ventricular hypertrophy, the chief effect is shortening of the QT interval and change in character of the ST segment (Fig 137).

QUINIDINE

Quinidine causes prolongation of all the electrical events of the heart. The QRS complex and QT interval are both prolonged by administration of quinidine. Most likely quinidine acts by altering the capability of the cell membrane to transmit excitation and recovery impulses. The direct action on the cell membrane decreases the threshold of irritability. This action enables quinidine to be effective in the treatment of arrhythmias.

By slowing the outward transmission of recovery along the cell membrane, endocardial recovery will create a force during the excited

state (ST segment) before the apparent stage of recovery (T wave). The ST segment change is similar to that seen in left ventricular hypertrophy. Quinidine may slow recovery to the extent it creates a large persistent everted cone inverting the T waves. Danger of cardiac stand still from quinidine is real if the ECG is not used to detect increasing QRS duration when large doses are used. With increasing concentration of quinidine, transmission of impulses is progressively slowed. The main ECG features of quinidine are:

- 1) Prolongation of PR interval
- 2) Increase in P wave size and duration
- 3) Increased QRS duration
- 4) ST segment depression
- 5) QT interval prolongation
- 6) T wave may become inverted

HYPOKALASSEMIA

When the concentration of potassium ions in the extracellular fluid decreases, there is an increased outward migration of potassium ions during the recovery phase. Recovery is closely related to the outward current of potassium ions. Apparently, when the intracellular and extracellular potassium ion concentration ratio reaches a critical level, the membrane recovers. With extracellular hypokalemia, a larger quantity of intracellular ions must migrate outward to create the critical ratio. The prolongation of the recovery phase enables the creation of wave fronts or currents within the endocardial shell. This is reflected in the ECG by ST segment changes.

The characteristic ECG findings in hypokalemia are:

- 1) Prolongation of the QT interval
- 2) ST segment deviations
- 3) Diminished amplitude of the T wave (Fig 138)

If hypokalemia persists, the constant increased outward migration of intracellular potassium ions results in cellular potassium depletion. Since the contractility and excitability of the cell is directly related to potassium ion

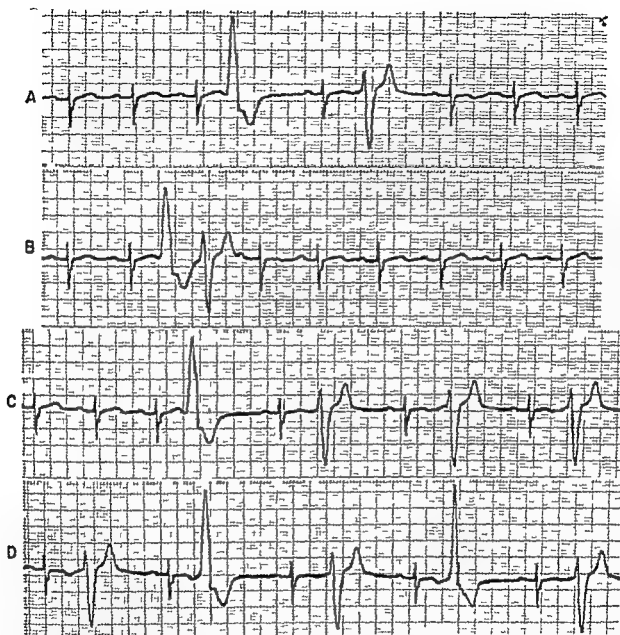


Figure 136 Digitalis intoxication. A) Ventricular premature contractions occurring from two foci. B) Coupling of impulses from the two foci. This may occur when the second focus initiates an impulse during the supranormal phase of excitability of the ventricle. C) The onset of bigeminy due to a ventricular premature every other beat. D) Alternating bidirectional bigeminy with the two foci alternating in creating ventricular premature contractions. All these rhythms occurred in a short time interval in one patient.

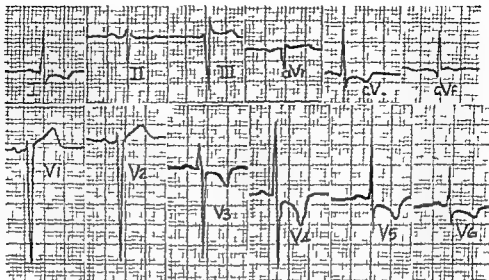


Figure 137 The same patient as seen in Fig 103 after digitalization. The QT interval is shortened from its previous duration.

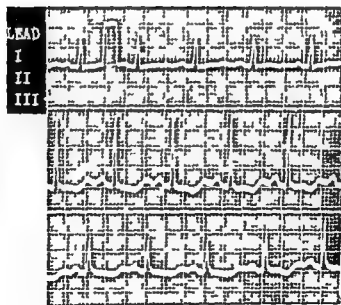


Figure 138 Prolongation of the QT interval and ST segment depression due to hypokalaemia.

concentration it is not surprising that arrhythmias occur.

Digitalis intoxication and hypokalaemia both create intracellular potassium ion depletion.

HYPERPOTASSAEMIA

Elevation of the extracellular potassium ion concentration shortens the recovery stage. A smaller amount of outward potassium ions migration is required to reach the critical intracellular/extracellular ratio. Recovery is rapidly transmitted to the epicardial surface increasing the demonstrable force of ventricular recovery. The shortening of the recovery period (ST interval) with increased force results in the tall peaked T waves of hyperkalaemia.

As the extracellular level of potassium ion concentration increases the outward driving force for potassium ion migration created by the difference in intracellular and extracellular ion concentration diminishes. When this force is effectively abolished standstill occurs. The atrium is first affected. P waves disappear due to atrial standstill. At progressively higher extracellular concentrations ventricular response is altered. Excitation becomes more difficult and the QRS complex widens. Fusion of the QRS and T waves occurs resulting in a scrambled electrocardiogram of biphasic waves. The distinction of excitation and recovery is not

possible. Ventricular standstill is the final phase of hyperkalaemia.

CALCIUM

Very little is known about the action of calcium ions in excitation and recovery. Diminished calcium ion concentration results in delay of the onset of recovery. This is manifested by prolongation of the QT interval with no apparent changes in character of the ST segment or T wave. Increased calcium ion concentration shortens the QT interval and may increase T wave amplitude or cause rounding of the T wave. Calcium has a synergistic action with digitalis.

THYRONIN

Thyrotoxicosis causes increased cardiac output. This is accomplished by increased heart rate with an increased or normal stroke volume. The ECG shows tachycardia and normal or increased QRS amplitude. Myxedema causes decreased cardiac output with diminished stroke volume and bradycardia. The ECG shows a slow heart rate and diminished QRS amplitude. The QRS amplitude may reflect either change in cell function with deposition of extracellular material in the heart or decreased stroke volume. Pericarditis with effusion may be an accompanying feature.

XIV

Arrhythmias

SINUS RHYTHMS

Normal Sinus Rhythm The SA node, atrium, AV node and ventricular endocardium are all under the influence of vagal inhibition and sympathetic acceleration. The SA node controls the cardiac rate and is called the pacemaker. In infants the normal rate may be between 100 and 150. The normal rate decreases with advancing age and in the adult the normal rate is 60 to 100.

When the impulse from the atrium reaches the AV node, there is a delay before ventricular excitation. This allows sufficient time for atrial systole prior to ventricular systole. In normal

in the SA node and traverses the rest of the cordia in normal fashion. The rate in the adult is over 100 per minute. Sinus tachycardia may be defined as a sinus rate above the normal rate for that particular individual. Sinus tachycardia is usually not thought to exceed a cardiac rate of 160 per minute. Rare instances of rates above that level have been noted. The onset of sinus tachycardia is insidious and its termination is gradual. The cardiac rate varies from minute to minute. Carotid sinus pressure causes perceptible smooth slowing of the cardiac rate. This is transient in nature and the original rate returns during pressure. Performing this maneuver during the recording of the tracing is



Figure 139 Normal sinus rhythm, rate 75 per minute

sinus rhythm the length of time for the impulse to pass from the SA node to the ventricle is dependent upon the length of time for the impulse to pass over the atria and the period of transmission through the AV conduction system. The normal adult PR interval is never less than .10 sec. and never more than .21 sec. In large adults with a cardiac rate below 70 the PR interval may be .21 sec. The method of exact measurement of the PR interval is discussed earlier in the text (Fig. 139).

Sinus Tachycardia The impulse originates

absolutely essential in establishing the diagnosis (Fig. 140).

Sinus tachycardia is occasionally associated with aberrant conduction. This permits an abnormal order of ventricular excitation resulting in abnormal QRS complexes.

Sinus tachycardia is due to changes in nervous regulation, myocardial function or increased venous return. With mild exertion the pulse rate increases 20 to 40 beats per minute but falls to normal limits within two minutes. When treatment is necessary, the first step is to correct

the underlying cause e.g. hyperthyroidism if such is present or replacement of blood in the face of acute blood loss

Sinus Bradycardia When a sinus mechanism is present at a rate of less than 60 per minute it is a sinus bradycardia. This may be a normal finding in young athletes. It has been reported present in individuals without demonstrable heart disease at rates as slow as 33 per minute. Sinus bradycardia with mild cardiac enlargement may be a normal finding and be merely

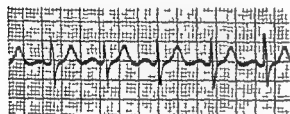


Figure 140 Sinus tachycardia rate 110 per minute

an athlete's heart with cardiac dilatation due to increased stroke volume

Sinus bradycardia is commonly seen in central nervous system involvement and certain infectious diseases including hepatitis. A relative bradycardia is seen in typhoid fever.

Sinus bradycardia is usually asymptomatic. Occasionally individuals will experience dizziness or syncope. It may be a manifestation of a sensitive carotid sinus and when associated with symptoms this reflex should be tested by massage of the carotid sinus during the time the ECG is recorded (Fig. 141).

Sinus Arrhythmia Sinus arrhythmia is the cyclic change in rate associated with respiration. The rate increases at the onset of inspiration and slows at the height of inspiration or at

the beginning of expiration. When this arrhythmia presents confusion in diagnosis having the patient hold his breath during the recording will disclose the nature of the disorder.

Vagal inhibition of the cardiac rate is at a minimum at the onset of inspiration (cross effect of the Hering-Breuer reflex). Increased venous return to the right heart occurs with inspiration due to negative intrathoracic pressure and the increased capacity of the pulmonary vascular bed.

When the breath is held in full inspiration there is often marked slowing of the heart rate. This reflex slowing may be more evident than carotid sinus slowing. I have observed momentary cardiac arrest and AV block from holding the breath in full inspiration.

Sinus arrhythmia is common in the young but it is also common in the elderly and the arteriosclerotic. It requires no treatment (Fig. 142).

Sinus Arrest In sinus arrest no impulse originates from the SA node and consequently the entire cardiac cycle is lost. The cycle which is lost may not equal a full length cycle. Sinus arrest cannot be clinically distinguished from sinoatrial block in which no impulse is transmitted from the SA node despite stimulation. If sinus arrest persists for several cycles without another rhythm occurring it is cardiac arrest and constitutes a cardiac emergency. Sinus arrest is usually precipitated by vagotonia and may be one manifestation of a hypersensitive carotid sinus (Fig. 143).

ATRIAL RHYTHM

Atrial Premature Contractions Whenever the electrical events of the heart are initiated

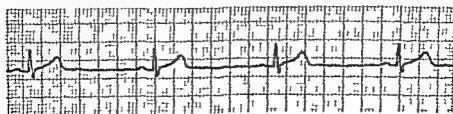


Figure 141 Sinus bradycardia rate 45 per minute



Figure 142 Sinus arrhythmia rate approximately 60 per minute

prematurely by a focus within the atria it is called an atrial premature contraction. The normal sequence of excitation does not occur and the P vector is altered. The PR interval may be entirely normal if the focus is far removed from the AV node. If the focus is close to the AV node the PR interval may become shortened.

The QRS complex is usually like those of normally conducted beats being associated with normal excitation. Less commonly early ventricular excitation occurs near the focus causing an abnormal QRS complex. This has been called aberrant conduction.

Usually there is a longer pause after the premature than the usual TP interval. This pause is not so long as to make a complete compensation for two successive cycles. This lack of compensatory pause is the chief method of establishing the diagnosis at the bedside. The pause is not compensatory because the atrial impulse causes the SA node to discharge thereby interrupting its normal rhythmic discharge. Whenever a premature from any focus fails to interrupt the normal rhythmic discharge of the SA

node and the node discharges during the prematurity the pause is completely compensatory.

When atrial premature contractions are noted in a patient with a history of previous rapid heart action it is a clue that the disorder is atrial tachycardia.

Atrial premature contractions 1) occur prematurely 2) are followed by a pause that is not completely compensatory 3) have a normal or shortened or even prolonged PR interval 4) have a normal or abnormally directed P vector and 5) commonly have a normal QRS complex (Fig 144).

Atrial Tachycardia An atrial premature contraction may initiate an attack of rapid heart action with an atrial focus as the pacemaker. This is atrial tachycardia. The rate is regular and is usually between 150 and 250 beats per minute. In infants it may be as rapid as 300 beats per minute.

The onset of the attack is sudden and it ends just as suddenly. A sudden motion may precipitate such an episode. The rate will not vary more than two beats per minute. When the episode is terminated there may be a fairly long



Figure 143 Sinus arrest

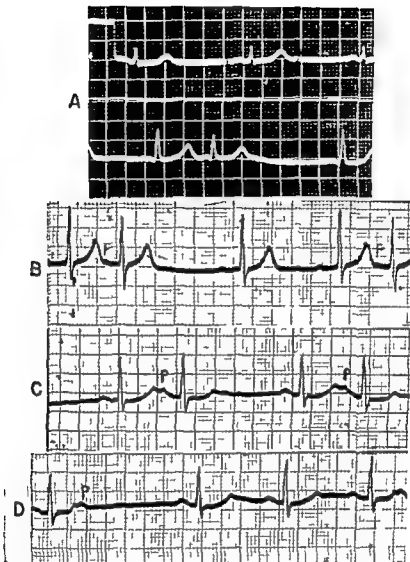


Figure 144 A) Normal rhythm above and premature occurring below B) Two atrial premature contractions C) Prematurities creating coupled rhythm D) A blocked atrial premature contraction

pause. Rarely has cardiac arrest occurred when the attack stopped. Short paroxysms of ventricular tachycardia have been said to occur at the time of termination.

Carotid sinus stimulation never affects the rate unless it stops the attack. This maneuver is not always successful and is of benefit in perhaps as few as 50% of the cases. Other manipulations to increase vagotonia may be used including eyeball pressure and gagging (Fig 145).

Atrial Flutter As the atria become more irritable they respond more rapidly to a single focus and their rate is increased. When atrial response to stimulation is as rapid as 200 to 360 per minute it is atrial flutter. The exact mechanism of atrial flutter is not known. It may be initiated with one focus or by multiple irritable foci. The electrocardiographic findings are the same and consist of a rapid atrial rate, a P vector directed in the axis of 90° and usually a regular ventricular response.

Only part of the atrial impulses are transmitted to the ventricle but this is commonly at a regular rate e.g. every other impulse transmitted or one in six impulses transmitted. The regular transmittal rate results in a regular ventricular response. If the atria are responding at a rate of 300 per minute and every other impulse is transmitted the ventricles will contract 150 times per minute. This is called a two to one

conduction associated with abnormal ventricular excitation (Fig 147)

NODAL RHYTHMS

Nodal Premature Contractions Any impulse originating from the AV node of necessity causes retrograde atrial excitation. This creates a P vector directed away from the AV node and toward the right shoulder. A PR interval of

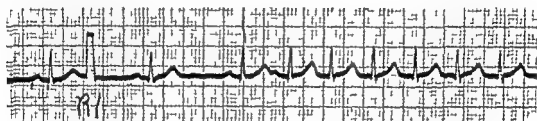


Figure 145 Paroxysmal atrial tachycardia

Flutter If only one in four impulses were transmitted the ventricular rate would be 75 per minute and this would be a four to one flutter. Ventricular response is not always regular and the flutter rate may vary between a three to one and a four to one rate. Many different combinations may exist. The regular atrial rate between 200 and 360 per minute with a regular ventricular response usually presents no difficulties in recognition (Fig 146)

Atrial Fibrillation When atrial excitation is as frequent as 350 to 500 per minute atrial fibrillation exists. The atria do not contract but merely fibrillate. The ventricular response is irregular with only a portion of the atrial impulses being transmitted to the ventricle. The ventricular rate is usually rapid in untreated cases. The ECG tracing will show small fibrillatory waves or no waves of atrial activity at all. The QRS complexes may show minor degrees of distortion

less than 10 seconds with a P vector compatible with a nodal focus is called a nodal impulse.

When an impulse originating from the node creates a premature contraction it is called a nodal premature contraction. These have been separated into three distinct groups: upper, middle, and lower nodal impulses. Upper nodal impulses have a nodal type P vector with the P wave just preceding the QRS complex. It is thought that an impulse originating high in the node will cause atrial excitation before the ventricle is stimulated. Middle nodal impulses are those with no visible P wave. The atrial events occur simultaneously with ventricular excitation and are lost. Lower nodal impulses allow the ventricle to be stimulated before atrial excitation. The QRS complex is followed by a P wave which behaves like a nodal type P wave (Fig 148)

Nodal premature contractions are followed by a com-

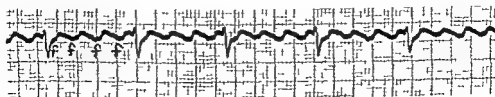


Figure 146 Atrial flutter is seen in Lead III

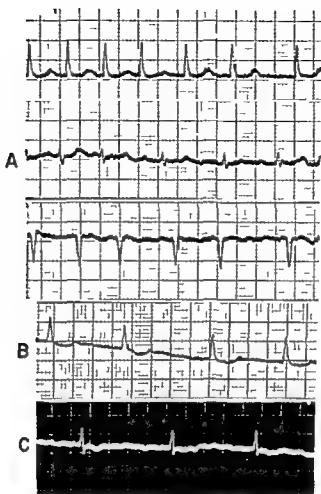


Figure 147 A) Rapid atrial fibrillation B) After digitalization C) So called flutter fibrillation

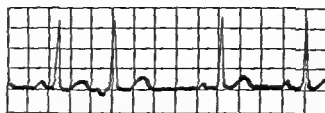
compensatory pause since the SA node discharges at its usual time and one SA impulse is lost (Fig 149). The QRS complex is usually normal. If the impulse originates low enough in the node it may be originating just from one main bundle and resemble a bundle branch block complex.

Nodal Rhythm When the pacemaker for the heart becomes the AV node, it is called nodal rhythm. The rate is slow, approximately 50 per minute, and regular. The AV node characteristically is slower than the SA node. The impulses look like the nodal impulses described above but are constant and comprise the basic rhythm. Nodal rhythm may be either upper, middle or lower nodal in origin (Fig 150).

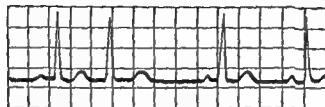
Nodal Tachycardia Whenever the AV node

emits impulses more rapidly than the SA node it becomes the cardiac pacemaker. When this rate exceeds 100 a minute, it is a nodal tachycardia. Like atrial tachycardia, it may occur in paroxysms with a sudden onset and sudden termination. Often the two conditions are clinically indistinguishable.

Nodal Escape If a period of sinus arrest occurs, a lower cardiac center usually takes over the job of pacemaker. The AV node is the next highest center and it may initiate the impulse. This is called nodal escape. The impulse has the usual characteristics of a nodal impulse. It differs from a nodal prematurity in that it follows a long pause or period of arrest and the prematurity occurs early. The impulse may de-



Upper Nodal Premature Contraction



Middle Nodal Premature Contraction



Lower Nodal Premature Contraction

Figure 148

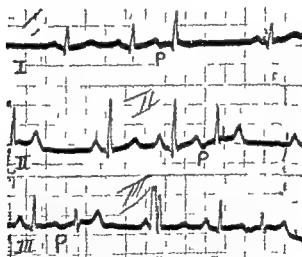


Figure 149 Nodal premature contractions

ascend further down the AV node to either bundle and finally descend to the ventricular muscle or the heart may cause excitation entirely. The gradual depression of the focus of origin is very important clinically, as it heralds the onset of cardiac arrest and is an emergency consideration.

Accelerated Nodal Conduction (See Chapter V)

VENTRICULAR RHYTHMS

Ventricular Prematurities When an impulse originates prematurely from the ventricle it is called a ventricular premature contraction. The impulse occurs early and is followed by a compensatory pause. The focus initiates ventricular excitation in only one ventricle. Thus the septum is activated only from one surface (Figs 151 and 152). The time for excitation is prolonged and the magnitude of the resultant vector is increased. This causes increased QRS

duration to 12 sec or more and marked increase in the size of the QRS complex.

Ventricular Tachycardia When tachycardia is the result of a ventricular pacemaker it is called ventricular tachycardia. The rate is usually between 150 and 250 per minute. Ventricular tachycardia is initiated by a ventricular prematurity and may be preceded by several prematurities just prior to onset. The rate may not be entirely regular and the cycle length may vary as much as 0.1 sec. The QRS complexes are widened to 12 sec or more. The P waves are slower and are at an independent rate.

The demonstration of the P waves is often absolutely necessary to establish a diagnosis due to the frequency of widened QRS complexes with those tachycardias having aberrant conduction (early ventricular excitation). Even a sinus tachycardia may be confused with ventricular tachycardia if the P waves are not

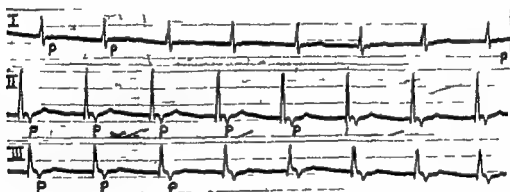


Figure 150 Nodal rhythm

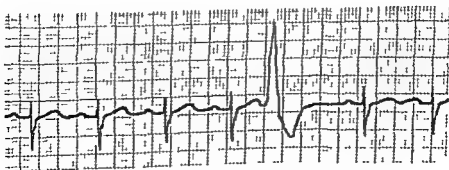


Figure 151 A simple ventricular premature contraction

found. Occasionally special leads including esophageal leads will be necessary to establish the presence of P waves.

Ventricular impulses from other foci may be interspersed throughout the record. Two foci may alternate, thus producing a bidirectional ventricular tachycardia. This is most often seen as a complication of digitalis intoxication. Termination of the tachycardia is followed by a full compensatory pause (Fig 153).

Ventricular Fibrillation Ventricular tachycardia may progress into ventricular fibrillation. The ventricles cease to contract and merely fibrillate at a rate between 130 and 500. The ventricular complexes may appear as fairly uniform oscillations of eight to ten millimeters

amplitude and then gradually decay with subsequent hypoxia. Terminally the oscillations appear as low amplitude undulations.

The atria continue to beat independently until hypoxia develops.

Recovery from ventricular fibrillation has been known to occur after a period of five minutes. It is nevertheless a serious emergency and is frequently death dealing.

AV BLOCK

First Degree AV Block When transmission through the AV conduction system is unduly prolonged it is called an AV block. If the delay results only in prolongation of the PR interval it is called a first degree AV block (Fig 154).

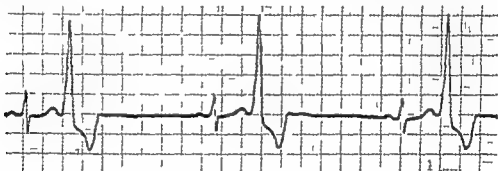


Figure 152 Ventricular premature contractions producing bigeminy

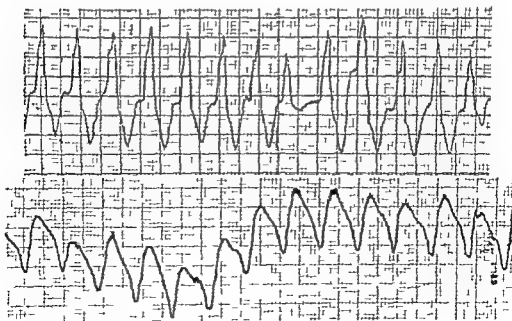


Figure 153 Two examples of ventricular tachycardia

Second Degree AV Block The delay may be great enough to prevent the transmission of the impulse part of the time. This results in a P wave without a following QRS complex. This is called a second degree AV block (Fig 155). When this occurs with progressive prolongation of the PR interval until a QRS complex is lost it is called a Wenckebach phenomenon (Fig 156).

Complete AV Block The inhibition at the AV node may be great enough to prevent the trans-

mission of atrial impulses altogether. In this instance the atria beat independent of the ventricles. This situation is called a third degree AV block or complete AV dissociation (Fig 157). The ventricular impulse may be initiated by a lower focus in the node in which case the ventricular complexes will appear normal or the impulse may originate in the ventricle with a wide QRS complex of a ventricular focus. In the latter instance the ventricular rate is commonly slower and it is called an idioventricular response.

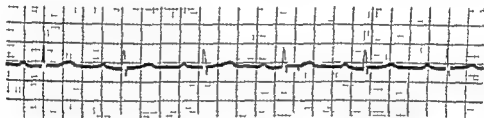


Figure 154 1st degree AV block



Figure 155 2nd degree AV block



Figure 156 Wenckebach's phenomenon

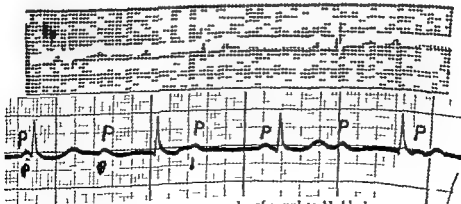


Figure 157 Two examples of complete AV block

XV

The Interpretation

THE ELECTROCARDIOGRAPHIC interpretation should be made in such a way as to give a maximum amount of information and a minimum amount of confusion. Dogmatic interpretations must be avoided as there are too many variables in the ECG. The maximum benefit of the tracing can only be obtained by clinical correlation.

The value of comparative records cannot be over emphasized. An inverted T3 may be a normal finding in one individual and an important change indicating disease in still another. Only adequate base line records will resolve the question on certain occasions.

The first part of the interpretation is entirely descriptive. One notes the basic rhythm e.g. normal sinus rhythm rate 75 per minute. Secondarily arrhythmias are noted such as normal sinus rhythm with occasional ventricular premature contractions. Any of the numerous arrhythmias may be present or combinations of arrhythmias e.g. complete AV block with sinus arrhythmia (sinus arrhythmia is noted in the P waves).

The intervals are described PR interval QRS duration QT interval. The character of the P waves may be described as normal increased amplitude or duration or abnormal characteristic notching or abnormally directed P vector. The QRS complex is described in terms of amplitude and configuration e.g. S1 S04 sec Q wave in Lead III and Lead II R R at V1. The ST segments are noted to be isoelectric elevated or depressed. The T wave configuration may be described as normal peaked increased amplitude or low amplitude. The presence of U waves is noted.

The mean QRS axis is determined in the

frontal plane in degrees and the transition noted in the V leads. This describes its spatial orientation. The T vector is described in axis degrees for the frontal plane and its transition in the V leads. From this information the spatial mean QRS T angle can be estimated from the spatial angle chart.

A simple descriptive interpretation is not much help to the clinician and requires no great skill. He wants to know whether the record is normal or abnormal and if possible what the abnormality means. This phase of the reading is the clinical interpretation. If the record is normal by the above measurements one should state *the record is within normal limits*. Any arrhythmia should be noted also in this portion of the reading.

If the PR interval was noted to be abnormally prolonged for the rate and age one states first degree AV block or other AV conduction disturbances are noted such as second degree AV block or complete AV block.

A prolonged QRS duration is interpreted in the clinical section as left bundle branch block right bundle branch block intraventricular conduction defect accelerated nodal conduction (WPW) S1 S0 pattern a normal variant or delay in activation over the right ventricle usually a normal variant. It is good to point out this in ominous sounding term (S1 S0 S1) is a normal variant and not worry the attending physician with trivial information.

If the QT interval was found to be abnormally prolonged it should be mentioned in the clinical interpretation as "prolonged QT interval consistent with _____". Electrolyte disturbance and rheumatic carditis are both

frequent causes. Often the attending physician will have asked for the tracing to detect changes consistent with some specific problem such as hypopotassemia.

Abnormally large P waves should be cited as consistent with atrial enlargement or dilatation. Don't try to distinguish between left and right atrial enlargement on the electrocardiogram alone. You will be wrong as often as right and it is better taste to give less information and be right when you give it.

When right axis deviation and precordial leads suggest its presence, one should say "right axis deviation consistent with right ventricular hypertrophy" or "right axis deviation with R R at V1 consistent with right ventricular enlargement or right bundle branch block."

As there are many causes for a wide split or mean QRS-T angle, it is probably best to note it as just an abnormality unless other changes accompany it. T wave changes after all can mean anything from myocardial infarction to changes secondary to a recent carbohydrate intake. Dogmatic interpretations without knowledge of the case do more harm than good. Serial electrocardiograms are very helpful in evaluating T wave abnormalities.

When a combination of findings suggesting left ventricular hypertrophy is present, it is probably safe to make the interpretation "abnormal record with increased QRS amplitude, widened QRS-T angle and ST segment changes consistent with left ventricular hypertrophy."

ST segment changes that do not appear normal should be noted, e.g., "ST segment changes consistent with epicardial injury." Suggest serial records for evaluation.

Classic infarctions may be diagnosed. Digitalis may be suspected. There are many valuable interpretations that can be made. One should never forget that an electrocardiogram does have limitations. It is used best with clinical information, previous comparative records and in some cases serial records. Because the electrocardiogram is often credited with infallibility by the uninitiated, avoid at all times dogmatic interpretations from the electrocardiogram alone.

There is still much valuable information to be learned from the use of the electrocardiogram. As yet we have only scratched the surface with its application. Even so, it is already a routine diagnostic procedure for both cardiac and general internal medicine problems.

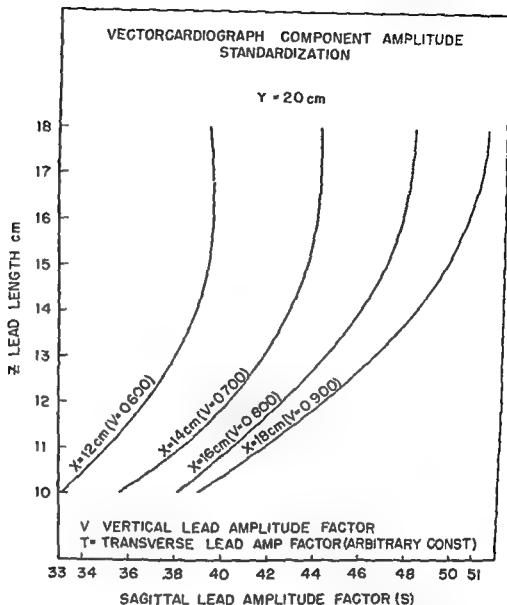


Figure 158 The amplification or standardization for the three vectorcardiographic leads (X , Y , Z) can be determined by 1) selecting the chart with the proper Y lead length 2) placing a straight edge at the left hand margin at the measured length of the Z lead 3) noting the point where the straight edge intersects the line representing the correct X lead length and 4) reading the amplification for Z (sagittal lead) directly beneath the point of intersection. The X amplification is given on the same curved line representing the X lead length. The amplification of Y is 1.00.

Given a Y lead of 20 cm length a Z lead of 17 cm length and an X lead of 12 cm length the amplifications are read as

$$\begin{aligned} Y &= 1.00 \\ X &= .60 \\ Z &= .40 \text{ (approximation)} \end{aligned}$$

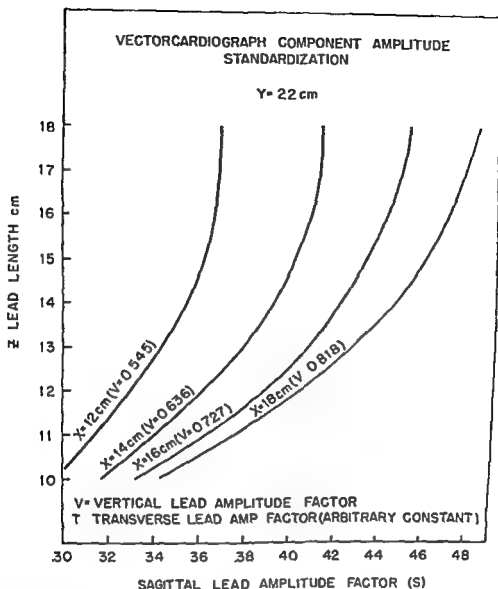


Figure 158 (Continued)

The most common lead lengths used in the adult are $Y = 30 \text{ cm}$, $T = 22 \text{ cm}$ and the sagittal varies. In such a case $Y = 1.00$, $T = 73$ and sagittal amplification is as follows:

Sagittal Lead	15 cm	=	36	amplification
	16 cm	=	38	
	17 cm	=	40	
	18 cm	=	41	
	19 cm	=	42	
	20 cm	=	43	
	21 cm	=	44	

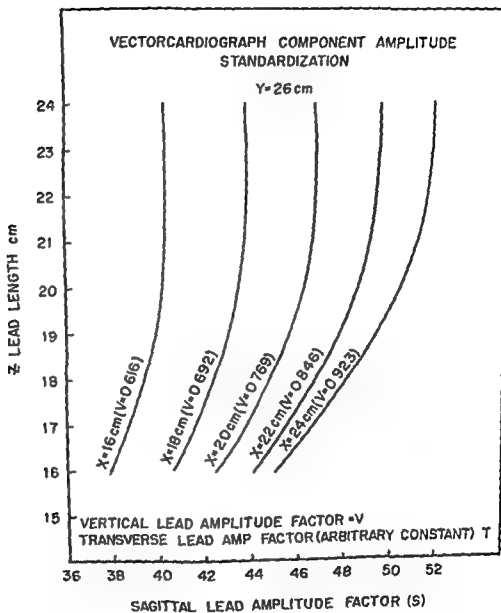


Figure 158 (Continued)

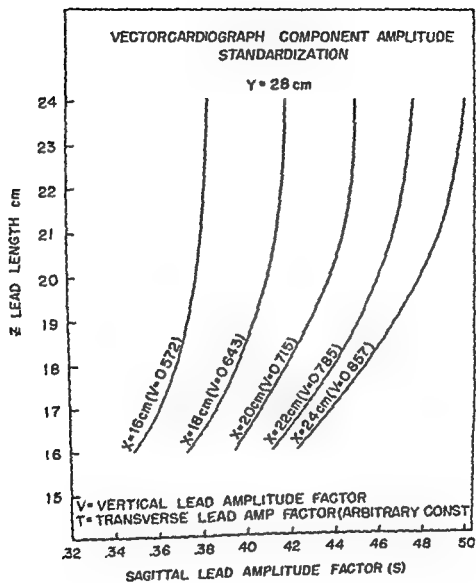


Figure 158 (Continued)

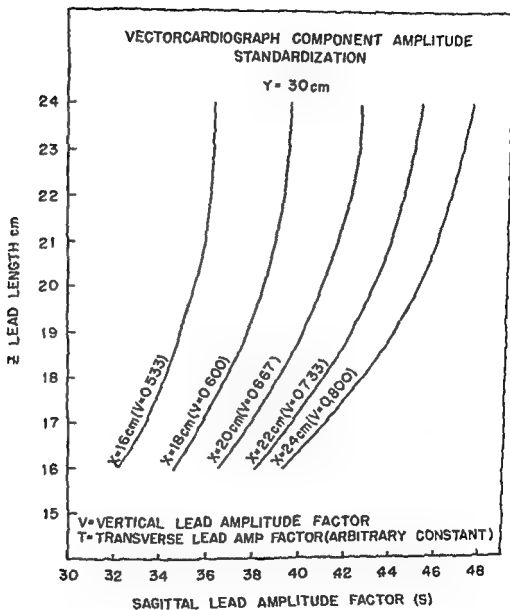


Figure 158 (Continued)

Index

A

- Aberrant conduction 122
- Absolute linear graph 111
- Accelerated conduction
 - See WPW
- Action potential of resting cell in membrane 14
- Atrial potential 26
- Atrial
 - anatomy of 19
 - enlargement 8
 - excitation of 19
 - excitation loop 19
- Arteriosclerotic heart disease 116
 - QRS amplitude in 116
 - QRS axis in 116
 - QRS T spatial angle in 116
 - QT interval in 116
 - ST segment in 116
 - T vector in 116
- Atrial fibrillation 126
 - atrial rate and 126
- Atrial flutter 125
 - axis of flutter (P) vector 125
 - flutter waves in 125
 - rate and 125
- Atrial premature contraction 124
 - compensatory pause in 124
 - paroxysmal rapid heart action and 124
- PR interval in 124
- Atrial tachycardia 124
 - cardiac rate in 124
 - sinus stimulation and 125
- Auricular fibrillation
 - See atrial fibrillation
- Auricular flutter
 - See atrial flutter
- Auricular premature contraction
 - See atrial premature contraction
- Auricular tachycardia
 - See atrial tachycardia
- A-V block
 - complete 131
 - first degree 130
 - second degree 131
- A-V node 20
- Axis deviation
 - left ventricular hypertrophy and 83-84
 - right ventricular hypertrophy and 86-88
- QRS 60

B

- Bazett's formula 56
- Bundle of Kent 98-99

C

- Calcium 121
 - digital and 121
 - QT interval and 121
 - T wave effects 121
- Cathodic ray oscilloscope 31
- Cell
 - excitation 14
 - injury 18
 - recovery 15
 - factors influencing 17
- Central terminal 36
- Compensatory pause 124
 - atrial premature contraction and 124
 - nodal premature contraction and 126
 - ventricular premature contraction and 129
- Conductor 27
 - body as a volume conductor 28
 - finite volume conductor 29
 - heterogeneous 27
 - homogeneous 27
 - infinite volume conductor 28
 - volume conductor 27-28
- Coordinate 3
- Coordinate graph 8
- Coronary artery experimental ligature 110-112

D

- Delay in activation over the right ventricle 13
- Delta wave 98-99
- Depolarization
 - See excitation
- Dielectric constant 14
- Digitalis 117
 - left ventricular hypertrophy and 117
 - normal action potential and 117
 - PR interval and 117
 - QT interval and 117
 - recovery 117
 - ST vector and 117
 - ST segment and 117
 - toxic manifestations 117
 - T vector and 117
- Dipole moment 29
- Doublet 12
- Doublet vector 14

F

- EKG instruments
 - direct writing 3
 - standards for operation 32
 - string gage galvanometer 30
- Einthoven's Law 34
- Einthoven's triangle 34
- Electric field 12-13
- Electric moment 29
- Electrode
 - exploring 36
 - negative 30
 - positive 30
- Excitation
 - atrial 19
 - cellular 14
 - ventricular 20
 - left bundle branch block and 91
 - polygon principle and 22-24
 - right bundle branch block and 93
 - summary of vector rules for 17
 - wave of 15
- Excited state 14
- Exploring electrode 36

F

- False bundle branch block
 - See WPW
- Fibrillatory waves 126
- Fleming's Law 30
- Frontal plane 5

G

- Galvanometer
 - string gage 30
- Gauss
 - direction of 16

H

- Haxial reference system 37
- Hypotassmia 118
 - QT interval and 118
 - ST segment and 118
 - T wave and 118
- Hypotassmia
 - clinical standard till and 111
 - I waves and 121
 - QRS complex and 121
 - T wave and 111

I

- Identification of maximum potential for the left ventricle 24

- in left ventricular hypertrophy 83 84
 - in physiological hypertrophy 83 84
 - in right ventricular hypertrophy 86 88
 - Index of potential seconds 71
 - Infarction
 - See myocardial infarction
 - Injury
 - cell 18
 - pericarditis and 105
 - Intraventricular conduction defect 96
 - Intrinsicoid deflection 49
 - Isopotential plane 13
- L**
- Law
 - Einthoven's 34
 - Fleming's 30
 - of cosines 65
 - of multiple simultaneity in vector addition 8
 - of parallelograms 6 33
 - of simple consecutive vector addition 7
 - of simple consecutive vector subtraction 7
 - Lead
 - augmented unipolar (aVR aVL aVF) 37
 - Einthoven's I II III 33
 - bipolar
 - sensitivity formula 33
 - length and vector axis 45
 - primal lead effect 39
 - positions for precordial leads 40
 - signal VCG 69
 - spatial reference system 42
 - standardization for ECG 31
 - transverse VCG 69
 - V 36
 - sensitivity 41
 - VCG reference system 68 71
 - VCG standardization 69
 - vertical VCG 69
 - Vr V1 V2 39
 - Left bundle branch block 91
 - QRS configuration in 91 92
 - QRS duration in 91
 - QRS loop in 91 92
 - recovery in 91 92
 - septal activation in 91 92
 - ST force in 91 92
 - T waves in 91 92
 - Left ventricular hypertrophy 82
 - axis deviation in 83 84
 - index of maximum potential in 83 84
 - QRS amplitude in 83
 - QRS duration in 83
 - QRS T spatial angle in 84
- QT interval in 84
 - potential seconds in 83 84
 - recovery in 83
 - ST segment depression in 83
 - T loop in 83 84
 - vertical el. circuit axis in 84
- M**
- Mean QRS axis 60
 - Mean vector 58
 - Membrane action potential
 - digitals and 117
 - resting cell and 14
 - reversal in excitation 14
 - Myocardial infarction 103 116
 - endocardial location 114
 - initial 01 sec QRS vector in 110 112 113
 - normal ECG in 109
 - Q wave in 110
 - QRS amplitude in 110
 - QRS notching in 115
 - ST segment in 110 112 113 115 116
 - ST vector in 110 112 113 115 116
 - T vector in 112
 - T waves in 109 110 112 113 116
 - Myocardium 121
 - pericarditis and 121
 - QRS amplitude in 121
- N**
- Negative electrode 30
 - Nodal escape 127
 - Nodal premature contraction 126
 - compensatory pause and 126
 - lower nodal 126
 - middle nodal 126
 - PR interval in 126
 - P vector and 126
 - retrograde conduction and 126
 - upper nodal 126
 - Nodal rhythm 127
 - Nodal tachycardia 127
 - Normal sinus rhythm 122
- O**
- Oersted experiment 15
 - Oscilloscope
 - cathode ray 31
- P**
- Pacemaker 19
 - Primal lead effect 39
 - and V leads 41
 - Paroxysmal tachycardia
 - in VFW 36 98
 - Pericarditis 105 107
 - myocardium and 121
 - myxodema and 121
 - QRS amplitude in 107
 - QRS loop in 105
 - recovery in 105 106
 - ST interval in 105
 - ST segment in 105 106
 - ST vector in 106
 - T vector in 106
- P**
- Plane 5
 - frontal 39
 - sagittal 44
 - transverse 40
 - P loop 47
 - P neutral 83
 - Potential 12
 - Potential seconds 24
 - for the left ventricle per cardiac cycle 24
 - left ventricular hypertrophy and 83 84
 - physiological hypertrophy and 84
 - Polygon
 - central resultant 6
 - principle 6
 - Positive electrode 30
 - Potassium current 15
 - P pulmonale 82
 - Precipitation
 - See VFW
 - PR interval 47
 - atrial premature contraction and 124
 - digitals and 147
 - first degree AV block and 130
 - quinniac and 118
 - nodal premature contraction and 126
 - VFW and 96 99
 - P vector
 - atrial flutter and 126
 - nodal premature contraction and 126
 - Wave 47
 - hyperpotassium and 121
 - ventricular tachycardia and 126
- Q**
- QRS amplitude 51
 - retrograde heart disease and 116
 - left ventricular hypertrophy and 83 84
 - myocardial infarction and 110
 - myxodema and 121
 - pericarditis and 107
 - right ventricular hypertrophy and 86 88
 - thyrotoxicosis and 121
 - ventricular premature contractions and 129

- QRS axis
arteriosclerotic heart disease and 116
normal range 60
- QRS complex
atrial premature contraction and 124
configuration 48
exercise and 54
quinidine and 118
respiration and 54
- QRS duration 51
hypertrophia and 121
left bundle branch block and 91
left ventricular hypertrophy and 83 84
right bundle branch block and 93
right ventricular hypertrophy and 86 88
S S S and 94
ventricular premature contraction and 129
ventricular tachycardia and 129
WPA and 96 99
- QRS loop
normal 72
in athletes 72
in children 72
in infants 72
in left ventricular hypertrophy 93
in left bundle branch block 91 92
in right ventricular hypertrophy 86 89
- QRS
mean vector 56 57
mean Q4 vector 57
mean Q4 vector of infarction 110 112 113
notching 49 115
- QRS T spiral angle 61
arteriosclerotic heart disease and 116
calculation 65
chart inside back cover
left ventricular hypertrophy and 83 84
right ventricular hypertrophy 86 89
- QT interval 56
arteriosclerotic heart disease and 116
calcium and 121
digitalis and 117
hypopotassemia and 118
left ventricular hypertrophy and 81
quinidine and 118
- Quinidine 118
cardiac standstill and 118
PR interval and 118
QRS complex and 118
QT interval and 118
- ST vector and 118
T waves and 118
- Q wave 48
of infarction 110
- R
Recovery
cellular 15
Factors influencing 17
digitalis and 117
left bundle branch block and 91 92
left ventricular hypertrophy and 83
pericarditis and 105 106
right bundle branch block and 93
summary of vector rules of 17
ventricular 24
wave of 15
- Remote electrode 29
- Repolarization
See recovery
- Resting cell 14
membrane reaction potential 14
- Retrograde conduction
nodal premature contraction and 126
- Right bundle branch block
excitation in 93
QRS configuration in 93
QRS duration in 93
recovery in 93
T wave in 93
- Right ventricular hypertrophy
anatomic changes in 84
axis deviation in 86 88
congenital heart disease and 89
index of minimum potential in 86 88
mitral stenosis and 89
QRS amplitude in 86 88
QRS configuration in 86 88
QRS duration in 86 88
QRS loop in 86 86
QRS T angle in 93 99
right bundle branch block and 93 95
- R wave 48
R wave 15
- S
Sagittal plane 7
Sinoauricular node 19
Sinus arrest 123
valsalva slowing with breath holding 123
Sinus bradycardia 123
athletes and 123
central nervous system disease and 123
stroke volume and 123
Sinus tachycardia 122
aberrant conduction and 122
carotid sinus pressure and 122
- Sodium current 14
- Spatial reference system 42
S S S
QRS configuration in 94
- Standardization
of ECG leads 31
of VEC leads 69
- Standards
for ECG instruments 32
- String gauge galvanometer 30
- ST segment 55
arteriosclerotic heart disease and 116
hypopotassemia and 115
left bundle branch block and 91 92
- ST segment
myocardial infarction and 110 112 113 115 118
pericarditis and 105
- ST vector
digitalis and 17
injury and 105
myocardial infarction and 110 112 113 115 118
pericarditis 106
quinidine and 118
- S wave 48
- T
Thyrotoxicosis
QRS and 121
tachycardia and 121
- Thyroxine 121
- Transverse plane 5
- Travel reference system 34
- T vector
arteriosclerotic heart disease and 116
digitalis and 117
myocardial infarction and 112
pericarditis and 106
- T wave
Atrial variability 55
calcium and 121
hypertrophia and 121
hypopotassemia and 118
left bundle branch block and 91 92
mean vector 60
myocardial infarction and 105 110 112 113 116
normal 55
quinidine and 118
right bundle branch block 93
WPA and 96 99
- U
U wave 26 56
- V
Vector
atrial excitation 19

- axis by inspection 37
 - axis and lead length 45
 - central resultants 6
 - components 4
 - coordinate value of 4
 - coordinate value calculation 5
 - definition 3
 - dipole 28
 - doublet 14
 - instantaneous 3
 - law of multiple simultaneous vector addition 7
 - law of simple consecutive vector addition 6
 - law of simple consecutive vector subtraction 7
 - lead axis rule 36
 - long left ventricular 23
 - loop of ventricular excitation 21 24
 - magnitude
 - instantaneous resultant vector versus polygon sides 11
 - time units and 10
 - mean 10 56
 - mean QRS 56 57 60
 - mean Q4 QRS 57 58
 - mean T 60
 - plane of 3
 - point of origin of 3
 - resultant 3
 - rule of excitation 15
 - rule of injury 18
 - rule of recovery 15
 - septal 22
 - ventricular recovery and 24
 - Ventricles
 - anatomy of 20
 - excitation of 20
 - polygon principle and 21 24
 - index of maximum potential of left 24
 - recovery of 24
 - Ventricular
 - fibrillation 130
 - hypertrophy
 - left 82
 - physiological 83
 - right 84
 - gradient 25
 - premature contractions 129
 - compensatory pause and 129
 - QRS duration and 127
 - recovery
 - cooling and warming and 24
 - endocardial delay in 24 26
 - vector loop of 25 26
 - septum 20
 - VCG
 - athletics 72
 - children's 72
 - frontal plane 66
 - infants 72
 - sagittal plane 66
 - leads 68 71
 - respiratory effects on 74
 - transverse plane 66
 - Ventricular tachycardia
 - P waves in 129
 - QRS duration and 129
- W
- Wave front 15
 - effective force of 18
 - ventricular excitation 20
 - ventricular recovery 24
 - Wenckebach phenomenon 131
 - Wilson's central terminal 36
 - Wolff Parkinson White syndrome
 - See WPW
- WPW
- paroxysmal tachycardia in 96
 - PR interval in 96 98
 - QRS duration in 96 99
 - T waves in 96 98
- Z
- Zero center 13 29
 - eccentric location 44

